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FILE COVERS 1907 - 21 Feb 2008 VOL 148 ISS 8 FILE LAST UPDATED: 20 Feb 2008 (20080220/ED)

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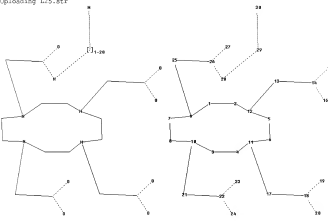
This file contains CAS Registry Numbers for easy and accurate substance identification. 'OBI' IS DEFAULT SEARCH FIELD FOR 'ZCAPLUS' FILE

=> d stat que L73 1.68 96 SEA FILE=ZCAPLUS ABB=ON PLU=ON GARLICH J?/AU L69 49 SEA FILE=ZCAPLUS ABB=ON PLU=ON SUHR R?/AU L70 710 SEA FILE=ZCAPLUS ABB=ON PLU=ON PATTERSON M?/AU 5 SEA FILE-ZCAPLUS ABB-ON PLU-ON L68 AND (L69 OR L70) L71 L72 4 SEA FILE-ZCAPLUS ABB-ON PLU-ON L69 AND L70 5 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L71 OR L72) L73

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=> d stat que L74
L25
              STR
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation: Uploading L25.str



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1 2 3 4 5 6 7 8 9 10 11 12
ring/chain nodes :
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13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 ring/chain bonds :

9-25 10-21 11-17 12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 22-24 22-23 25-26 26-28 26-27 28-29 29-30 ring bonds :

ring nodes :

1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10

exact/norm bonds :

1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10 9-25 10-21 11-

12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 22-24 22-23 25-26 26-28

26-27 28-29

29-30

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS

22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS

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=> s L73 L76 1 L73

=> file wpix

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FILE LAST UPDATED: 20 FEB 2008 <20080220/UP>
MOST RECENT THOMSON SCIENTIFIC UPDATE: 200812 <200812/DW>
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>>> IPC Reform backfile reclassification has been loaded to the end of November 2007. No update date (UP) has been created for the reclassified documents, but they can be identified by 20060101/UPIC and 20061231/UPIC, 20070601/UPIC, 20071001/UPIC and 20071130/UPIC. <<</p>

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FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE http://scientific.thomson.com/support/patents/coverage/latestupdates/

EXPLORE DERWENT WORLD PATENTS INDEX IN STN ANAVIST, VERSION 2.0: http://www.stn-international.com/archive/presentations/DWPIAnaVist2_0710.pdf

- >>> XML document distribution format now available See HELP XMLDOC <<<
- >>> ECLA Codes and Current US National Classifications have been added see NEWS and HELP CHANGE <<<
- >>> HELP for European Patent Classifications see HELP ECLA, HELP ICO <<<

>>> Updated PDF files in the following links: http://www.stn-international.de/stndatabases/details/ico_0801.zip http://www.stn-international.de/stndatabases/details/epc 0801.zip <<< 'BIX' IS DEFAULT SEARCH FIELD FOR 'WPIX' FILE => s L73 34 GARLICH J?/AU

L77

32 SUHR R?/AU 178 PATTERSON M?/AU 32 SUHR R?/AU 178 PATTERSON M?/AU 2 (L71 OR L72)

=> file stnguide

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FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: Feb 15, 2008 (20080215/UP).

=> dup rem L75 L76 L77 FILE 'ZCAPLUS' ENTERED AT 10:22:04 ON 21 FEB 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

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L78 5 DUP REM L75 L76 L77 (3 DUPLICATES REMOVED) ANSWERS '1-5' FROM FILE ZCAPLUS

=> d ibib abs hitind hitstr L78 1-5

L78 ANSWER 1 OF 5 ZCAPLUS COPYRIGHT 2008 ACS on STN DUPLICATE 1

ACCESSION NUMBER: 2005:324033 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 142:379479

TITLE: Chelate based scaffolds in tumor targeting INVENTOR(S): Garlich, Joseph R.; Suhr, Robert G.; Patterson, Mary

PATENT ASSIGNEE(S): Semafore Pharmaceuticals Inc., USA

SOURCE: PCT Int. Appl., 47 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Pat.ent.

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	PATENT NO.					D	DATE		APPLICATION NO.							DATE			
					_														
WO	WO 2005032599				A1 20050414			WO 2004-US32289						20040930					
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,		
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,		
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,		

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NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
            TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
        RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
            AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
            EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
            SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
            SN. TD. TG
    EP 1684809
                               20060802
                                          EP 2004-789423
                         A1
                                                                  20040930
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK
    US 2007104645
                        A1
                              20070510
                                           US 2006-573938
                                                                  20060725
PRIORITY APPLN. INFO.:
                                                               P 20030930
                                           US 2003-507427P
                                           WO 2004-US32289
                                                               W 20040930
     This invention relates to novel complexes that can be used to target tumor
     cells. The complexes include a ligand including a tetraazacyclododecane
     macrocycle ring core that can bind metal ions including radioactive lanthanide
     ions. The complexes can mimic ανβ3 integrin receptor antagonists and deliver
     the complexed radioactive metals to the tumor cells. For example, 24.4 mM of
     cyclen reacted with 24.4 mM of tert-Bu bromoacetate to give 5.72 g 1.4-DO2A
     bis-tert-Bu ester (82% of theory) as clear viscous oil. The oil was dissolved
     in MeOH, allowed to crystallize, the solid obtained was filtered, washed with
     water and then dried to give 4.3964 g of white solid.
    ICM A61K051-00
    63-8 (Pharmaceuticals)
    Section cross-reference(s): 1, 8, 24
    849610-60-2P 849610-61-3P 849610-62-4P 849610-63-5P
    849610-64-6P 849610-65-7P 849610-66-8P
    849610-67-9P 849610-68-0P 849610-69-1P
    849610-70-4P 849610-71-5P 849610-72-6P
    849610-73-7P 849610-74-8P 849610-75-9P
    849610-76-0P 849610-77-1P 849610-78-2P
    849610-79-3P 849610-80-6P 849610-81-7P
    849610-82-8F 849610-83-9P 849610-84-0P
    849610-85-1P 849610-86-2P 849610-87-3P
    849610-88-4P 849610-89-5P 849610-90-8P
    849610-91-9P 849610-92-0P 849610-93-1P
    849610-94-2P 849610-95-3P 849610-96-4P
    849610-97-5P
                  849610-98-6P 849610-99-7P 849611-00-3P
    849680-38-2P
    RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
    study); PREP (Preparation); USES (Uses)
        (chelate-based scaffolds for tumor targeting)
    849610-60-2P 849610-65-7P 849610-66-8P
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849610-70-4P 849610-71-5P 849610-72-6P
849610-73-7P 849610-74-8P 849610-75-9P
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849610-82-8P 849610-83-9P 849610-84-0P
849610-85-1P 849610-86-2P 849610-87-3P
849610-88-4P 849610-89-5P 849610-90-8P
849610-91-9P 849610-92-0P 849610-93-1P
949610-94-2P 849610-95-3P 849610-96-4P
849680-88-29
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RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(chelate-based scaffolds for tumor targeting)

849610-60-2 ZCAPLUS RN

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, $\alpha 4-[[(4$ acetylphenyl)amino[carbonyl]-10-[2-[(3-aminopropyl)amino]-2-oxoethyl]-, α4-ethyl ester (9CI) (CA INDEX NAME)

- RN 849610-65-7 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminohexyl)amino]-2-oxoethyl]-α-[2-[(2-carboxybenzoyl)amino]ethyl]-(9cI) (CA INDEX NAME)

- RN 849610-66-8 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(5-aminopentyl)amino]-2-oxoethyl]-α-[2-((2-carboxybenzoyl)amino]ethyl]-(9CI) (CA INBEX NAME)

RN 849610-67-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-aminobutyl)amino]-2-oxoethyl]- α -[2-[(2-carboxybenzoyl)amino]ethyl]-(CA INDEX NAME)

RN 849610-68-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(3-aminopropyl)amino]-2-oxoethyl]-a-(carboxymethyl)- (9CI) (CA INDEX NAME)

- RN 849610-69-1 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-aminobutyl)amino]-2-oxoethyl]- α -(carboxymethyl)- (9CI) (CA INDEX NAME)

- RN 849610-70-4 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(5-aminopentyl)amino]-2-oxoethyl]-a-(carboxymethyl)- (9CI) (CA INDEX NAME)

- RN 849610-71-5 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminohexyl)amino]-2-oxoethyl]- α -(carboxymethyl)- (9C1) (CA INDEX NAME)

- RN 849610-72-6 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, $10-[2-[[3-[(aminoiminomethy1)amino]propy1]amino]-2-oxoethy1]-<math>\alpha$ -(carboxymethy1)-(9C1) (CA INDEX NAME)

- RN 849610-73-7 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[4-[(aminoiminomethyl)amino]butyl]amino]-2-oxoethyl]-a-(carboxymethyl)-(9C1) (CA INDEX NAME)

- RN 849610-74-8 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[5-[(aminoiminomethy1)amino]penty1]amino]-2-oxoethy1]-a-(carboxymethy1)-(9C1) (CA INDEX NAME)

- RN 849610-75-9 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[(aminoiminomethyl)amino]hexyl]amino]-2-oxoethyl]- α -(carboxymethyl)-(9CI) (CA INDEX NAME)

RN 849610-76-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]-α-(2-carboxyethyl)- (CA INDEX NAME)

RN 849610-77-1 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(3-aminopropyl)amino]-2-oxoethyl]- α -(2-carboxyethyl)- (9CI) (CA INDEX NAME)

RN 849610-78-2 ZCAPLUS

CN 1,4,7,10-Tetrazzacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-aminobutyl)amino]-2-oxoethyl]- α -(2-carboxyethyl)- (9CI) (CA INDEX NAME)

- RN 849610-79-3 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(5-aminopentyl)amino]-2-oxoethyl]-α-(2-carboxyethyl)- (9CI) (CA INDEX NAME)

- RN 849610-80-6 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminohexyl)amino]-2-oxoethyl]-a-(2-carboxyethyl)- (9CI) (CA INDEX NAME)

- RN 849610-81-7 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]- α -(carboxymethyl)- (9CI) (CA INDEX NAME)

- RN 849610-82-8 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, $10-[2-[[2-(aminoiminomethy1)amino]ethy1]amino]-2-oxoethy1]-<math>\alpha$ -(2-carboxyethy1)-(CA INDEX NAME)

RN 849610-83-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[(aminoimnomethyl)amino]propyl]amino]-2-oxoethyl]- α -(2-carboxyethyl)- (CA INDEX NAME)

RN 849610-84-0 ZCAPLUS

RN 849610-85-1 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[5-[(aminoiminomethyl)amino]pentyl]amino]-2-oxoethyl]-α1-(2carboxyethyl)- (9CI) (CA INDEX NAME)

RN 849610-86-2 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[aminoiminomethyl]amino]hexyl]amino]-2-oxoethyl]- α 1-(2-carboxyethyl)-(9C1) (CA INDEX NAME)

RN 849610-87-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, $10-[2-[\{2-(\{aminoiminomethyl\}amino]ethyl]amino]-2-oxoethyl]-<math>\alpha$ 1-(carboxymethyl)-(9C1) (CA INDEX NAME)

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RN 849610-88-4 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]- α l-(3-carboxypropyl)- (9CI) (CA INDEX NAME)

- RN 849610-89-5 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, $10-[2-[(3-\min opropy1) \min o]-2-oxoethy1]-\alpha1-(3-carboxypropy1)- (9CI) (CA INDEX NAME)$

- RN 849610-90-8 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-aminobutyl)amino]-2-oxoethyl]-αl-(3-carboxypropyl)- (9CI) (CA INDEX NAME)

- RN 849610-91-9 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(5-aminopenty1)amino]-2-oxoethy1]- α 1-(3-carboxypropy1)- (CA INDEX NAME)

RN 849610-92-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminohexyl)amino]-2-oxoethyl]-α1-(3-carboxypropyl)- (9CI) (CA INDEX NAME)

RN 849610-93-1 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[(aminoiminomethyl) amino]propyl]amino]-2-oxoethyl]-α1-(3-carboxypropyl)- (CA INDEX NAME)

RN 849610-94-2 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[4-[aminoiminomethyl)amino]butyl]amino]-2-oxoethyl]-al-(3carboxypropyl)- (CA INDEX NAME)

RN 849610-95-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[5-[aminoiminomethyl)amino]pentyl]amino]-2-oxoethyl]- α 1-(3-carboxypropyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{CO2H} \\ \text{HO2C-CH2} \\ \text{III} \\ \text{CH2-CO2H} \\ \text{CH2-CO2H} \\ \text{CH2-CO2H} \\ \text{CH2-CO2H} \\ \text{CH3-CO2H} \\ \text{CH3-CO3H} \\ \text{CH$$

RN 849610-96-4 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[(aminoiminomethyl) amino]hexyl]amino]-2-oxoethyl]-α1-(3-carboxypropyl)- (9CI) (CA INDEX NAME)

RN 849680-88-2 ZCAPLUS

CN Yttrate(1-), [10-[2-[[5-[(aminoiminomethyl)amino]pentyl]amino]-2-(oxo-KO)ethyl]-d1-(carboxymethyl)-1, 4, 7, 10-tetraazacyclododecane-1, 4, 7-triacetato(5-)-KN1, KN4, kN7, KN10, KO1, .k appa.04, KO7]-, hydrogen (9CI) (CA INDEX NAME)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L78 ANSWER 2 OF 5 ZCAPLUS COPYRIGHT 2008 ACS on STN DUPLICATE 2 2004:878386 ZCAPLUS Full-text

ACCESSION NUMBER:

DOCUMENT NUMBER: 141:366126 TITLE:

Preparation of quaternized derivatives of (morpholinyl)phenylbenzopyranone as Pi-3 kinase

inhibitor prodrugs

INVENTOR(S): Garlich, Joseph R.; Durden, Donald L.; Patterson, Mary; Su, Jingdong; Suhr, Robert G.

PATENT ASSIGNEE(S): Semafore Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 136 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent.

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAI	ENT I				KIND DATE			APPLICATION NO.							DATE			
WO 2004089925					A1		20041021			WO 2004-US10399 200								
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		ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	
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		TD,	TG															
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GI

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The invention provides methods to prepare prodrugs I [Z and Z1-3 independently AB = 0 or S; R1 and R2 independently = H, (un)substituted- aliphatic, -aryl, OH, CN, halo, etc.; R3 = H, (un)substituted-aliphatic, -aryl; R4 and R5 = H, (un)substituted-aliphatic, -aryl, heterocyclyl, aryloxy, carboxy, or taken together form an (un) substituted heterocycle; R6 = H, (un) substitutedaliphatic, -aryl, etc.; R7 = -CH2-, -CH(CH3)-, -CH(Ph)-, -C(CH3)(CO2H)- or CH(CH(CH3)2)-; T is optional but when present = targeting agent], possessing a hydrolyzable quaternary nitrogen which can provide metabolites II capable of inhibiting PI-3 kinase. Thus, e.g., III was prepared via N-alkylation of IV with chloromethyl-t-butylsuccinate followed by hydrolysis and chlorination to the acid chloride which was reacted with resin bound peptide (arg-gly-asp-ser) after which cleavage from the resin provided III. III was evaluated for in vivo efficacy against non-small cell lung cancer and after 17 days a 35% reduction in tumor volume was observed (at 25mg/kg/day dosage). The novel compds. are IV and analogs thereof comprising a reversibly quaternized amine.
- ICM C07D311-22
- ICS C07D407-12; C07D475-04; A61K031-5377; A61P025-00 27-14 (Heterocyclic Compounds (One Hetero Atom))
- Section cross-reference(s): 1, 34, 63

REFERENCE COUNT: THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L78 ANSWER 3 OF 5 ZCAPLUS COPYRIGHT 2008 ACS on STN DUPLICATE 3

2004:891169 ZCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 142:322489

TITLE: Nanoparticles for delivery of pifithrins to combat cell death due to chemotherapy and radiation Brannon-Peppas, L.; Soehl, K.; Monaco, M. D.; AUTHOR(S):

Garlich, J.; Patterson, M.; Smith, T. C. Department of Biomedical Engineering and Division of CORPORATE SOURCE:

Pharmaceutics, The University of Texas at Austin,

Austin, TX, 78712-0231, USA

SOURCE: Journal of Drug Delivery Science and Technology

(2004), 14(4), 257-264 CODEN: JDDSAL

PUBLISHER: Editions de Sante

DOCUMENT TYPE: Journal LANGUAGE: English

This work describes the first stage of our research efforts to develop targetable nanoparticles to deliver agents to help healthy bone marrow cells survive radiation and chemotherapy. Administering pifithrin, a small mol. inhibitor of the protein p53, could prevent p53 initiated cell death. The p53

protein imparts sensitivity to normal tissue subjected to genotoxic stress such as radiation therapy or chemotherapy. We describe the conversion of pifithrin- α to pifithrin- β in buffer and serum and even while frozen and the implications in developing successful formulations. Encapsulation of pifithrin-β in biodegradable nanoparticles of poly(lactic-co-glycolic) acid showed encapsulation of up to 13% pifithrin and release in vitro of at least 28 days. Particle sizes ranged from 240 to 3250 nm, depending on the preparation methods used including variation of organic solvent type and amount

63-5 (Pharmaceuticals)

Section cross-reference(s): 1, 8

REFERENCE COUNT: THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L78 ANSWER 4 OF 5 ZCAPLUS COPYRIGHT 2008 ACS on STN 2004:435190 ZCAPLUS Full-text ACCESSION NUMBER:

TITLE: Targeted Delivery of p53 Inhibitors

Smith, Tim C.; Garlich, Joseph R.; Patterson, Mary AUTHOR(S):

L.: Suhr. Robert G.

CORPORATE SOURCE: Semafore Pharmaceuticals, Indianapolis, IN, 46268, USA SOURCE: Abstracts, 36th Central Regional Meeting of the

American Chemical Society, Indianapolis, IN, United States, June 2-4 (2004), GEN-452. American Chemical

Society: Washington, D. C. CODEN: 69FMAU

DOCUMENT TYPE: Conference; Meeting Abstract

The protein p53 is a tumor suppressor, which often is triggered during chemoand radiation therapy, causing unwanted side effects by inducing apoptosis of healthy tissue such as the hematopoietic system. Thus suppression of p53 in healthy tissues during therapy should decrease the damage. Pifithrin- and pifithrin- have been shown to act as small mol. inhibitors of p53. We have embarked on a program to target pifithrin- and to bone, thus offering selective protection to bone marrow and the immune system during therapy. This presentation will focus on the synthetic chemical of linking bone-seeking moieties to pifithrin- and as well as promising preliminary in vitro studies.

L78 ANSWER 5 OF 5 ZCAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:435232 ZCAPLUS Full-text

TITLE: Novel Purification Techniques and the Solid Phase

Synthesis of Macrocyclic Ligands AUTHOR(S):

Garlich, Joseph R.; Patterson, Mary; Smith, Tim

C.; Suhr, Robert G.; Georgiadis, Taxiarchis M.

Semafore Pharmaceuticals, Indianapolis, IN, 46268, USA

SOURCE: Abstracts, 36th Central Regional Meeting of the American Chemical Society, Indianapolis, IN, United

States, June 2-4 (2004), INV-033. American Chemical

Society: Washington, D. C.

CODEN: 69FMAU

CORPORATE SOURCE:

DOCUMENT TYPE: Conference: Meeting Abstract

One highly useful procedure in parallel or combinatorial synthesis is the clean-up of reaction mixts. using facilitated liquid-liquid extraction Researchers have previously described the use of large mesh sized diatomaceous earth beads coated with an aqueous phase for simultaneous extraction workup of an array of compds. simply by exposure of the reaction mix dissolved in an organic phase to the beads. We have taken this concept beyond simple liquidliquid extns. by employing diatomaceous earth beads coated with various aqueous based scavenging, catalytic and reactive solns. This supported aqueous film exposure can be utilized during a reaction to introduce catalysts or reactive reagents which react at the films water-organic interface. Post-

reaction workup is thus reduced to simple filtration or decanting. Novel phys. formats for this technique have also been explored. This work and the solid-phase synthesis of macrocyclic ligands will be discussed.

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STRUCTURE FILE UPDATES: 20 FEB 2008 HIGHEST RN 1004854-20-9 DICTIONARY FILE UPDATES: 20 FEB 2008 HIGHEST RN 1004854-20-9

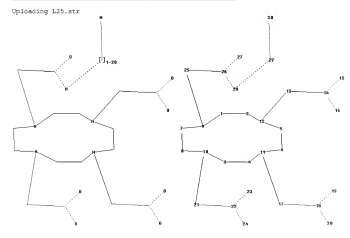
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http://www.cas.org/support/stngen/stndoc/properties.html



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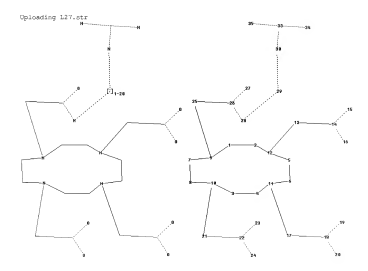
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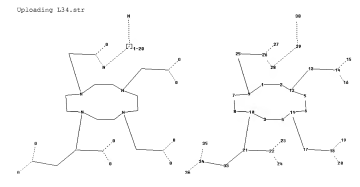
10/573938 ring nodes :

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1 2 3 4 5 6 7 8 9 10 11 12
ring/chain nodes :
13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 33 34 35
ring/chain bonds :
9-25 10-21 11-17 12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 22-24
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22-23 25-26 26-28 26-27 28-29 29-30 30-33 33-34 33-35 ring bonds : 1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10 exact/norm bonds :

1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10 9-25 10-21 11-17 12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 22-24 22-23 25-26 26-28 26-27 28-29 29-30 30-33 33-34 33-35

Match level : 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 33:CLASS 34:CLASS 35:CLASS



ring nodes : 1 2 3 4 5 6 7 8 9 10 11 12 ring/chain nodes : 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 33 34 35 ring/chain bonds :

9-25 10-21 11-17 12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 21-33

```
ring bonds:
1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10
exact/norm bonds:
1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10 9-25 10-21 11-17
17
18-20 18-19 18-19 18-20 18-19 21-22 21-33 22-24 22-23 25-26
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22-24 22-23 25-26 26-28 26-27 28-29 29-30 33-34 34-35 34-36

26-28 26-27 28-29 29-30 33-34 34-35 34-36

Match level :

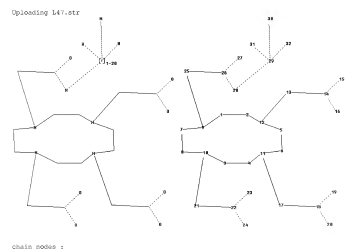
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33:CLASS 34:CLASS 35:CLASS 36:CLASS



31 32

31 32 ring no

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12

ring/chain nodes:

29-31 29-32

ring/chain bonds :

9-25 10-21 11-17 12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 22-24 22-23 25-26 26-28 26-27 28-29 29-30

ring bonds :

1-2 1-9 2-12 3-4 3-10 4-11 5-6 5-12 6-11 7-8 7-9 8-10

exact/norm bonds :

 $1-2 \quad 1-9 \quad 2-12 \quad 3-4 \quad 3-10 \quad 4-11 \quad 5-6 \quad 5-12 \quad 6-11 \quad 7-8 \quad 7-9 \quad 8-10 \quad 9-25 \quad 10-21 \quad 11-11-11 \quad 11-11-11-11 \quad 11-11-11 \quad 11-11-11 \quad 11-11-11 \quad 11-11-11 \quad 11-11-11 \quad 1$

12-13 13-14 14-16 14-15 17-18 18-20 18-19 21-22 22-24 22-23 25-26 26-28 26-27 28-29

29-30 29-31 29-32

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS

19:CLASS 20:CLASS 21:CLASS

22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS

31:CLASS 32:CLASS

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=> d stat que L32 L25 STR

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Structure attributes must be viewed using STN Express query preparation. L27 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

L29 2020 SEA FILE=REGISTRY SSS FUL L25

L31 62 SEA FILE=REGISTRY SUB=L29 SSS FUL L27

L32 9 SEA FILE=ZCAPLUS ABB=ON PLU=ON L31

=> d stat que L37 L25 STF

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express guery preparation.

L29 2020 SEA FILE=REGISTRY SSS FUL L25

L34 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation. L36 12 SEA FILE=REGISTRY SUB=L29 SSS FUL L34

L36 12 SEA FILE=REGISTRY SUB=L29 SSS FUL L34
L37 1 SEA FILE=ZCAPLUS ABB=ON PLU=ON L36

=> d stat que L45

L2

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-3/BI OR 849680-88-2/BI OR 95196-95-5/BI) L25 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

L29 2020 SEA FILE=REGISTRY SSS FUL L25

L31 62 SEA FILE=REGISTRY SUB=L29 SSS FUL L27

L32 9 SEA FILE-ZCAPLUS ABB-ON PLU-ON L31

L34 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

10/573938	
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L42	75 SEA FILE=REGISTRY ABB=ON PLU=ON L40 AND M/ELS
L45	8 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L41 OR L42) AND L38
	· · · · · · · · · · · · · · · · · · ·
-> 0 0+0+	mus TEE

=> d stat que L55 L25

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Structure attributes must be viewed using STN Express query preparation. L29 2020 SEA FILE=REGISTRY SSS FUL L25

L47 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express guery preparation.

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L50 142 SEA FILE=REGISTRY ABB=ON PLU=ON L49 AND M/ELS L54 9 SEA FILE=REGISTRY ABB=ON PLU=ON L50 AND Y/ELS

10 SEA FILE=ZCAPLUS ABB=ON PLU=ON L54 L55

=> d stat que L67

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OR 7440-10-0/BI OR 7440-12-2/BI OR 7440-19-9/BI OR 7440-20-2/BI OR 7440-27-9/BI OR 7440-30-4/BI OR 7440-45-1/BI OR 7440-52-0/B I OR 7440-53-1/BI OR 7440-54-2/BI OR 7440-60-0/BI OR 7440-64-4/ BI OR 7440-65-5/BI OR 849610-60-2/BI OR 849610-61-3/BI OR 849610-62-4/BI OR 849610-63-5/BI OR 849610-64-6/BI OR 849610-65 -7/BI OR 849610-66-8/BI OR 849610-67-9/BI OR 849610-68-0/BI OR 849610-69-1/BI OR 849610-70-4/BI OR 849610-71-5/BI OR 849610-72 -6/BI OR 849610-73-7/BI OR 849610-74-8/BI OR 849610-75-9/BI OR 849610-76-0/BI OR 849610-77-1/BI OR 849610-78-2/BI OR 849610-79 -3/BI OR 849610-80-6/BI OR 849610-81-7/BI OR 849610-82-8/BI OR 849610-83-9/BI OR 849610-84-0/BI OR 849610-85-1/BI OR 849610-86 -2/BI OR 849610-87-3/BI OR 849610-88-4/BI OR 849610-89-5/BI OR 849610-90-8/BI OR 849610-91-9/BI OR 849610-92-0/BI OR 849610-93 -1/BI OR 849610-94-2/BI OR 849610-95-3/BI OR 849610-96-4/BI OR 849610-97-5/BI OR 849610-98-6/BI OR 849610-99-7/BI OR 849611-00 -3/BI OR 849680-88-2/BI OR 95196-95-5/BI)

1.25 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation. L27 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

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L32 9 SEA FILE=ZCAPLUS ABB=ON PLU=ON L31

L34 STR

L40

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

L36 12 SEA FILE=REGISTRY SUB=L29 SSS FUL L34 L37 1 SEA FILE=ZCAPLUS ABB=ON PLU=ON L36

L38 9 SEA FILE=ZCAPLUS ABB=ON PLU=ON L37 OR L32

273 SEA FILE=REGISTRY ABB=ON PLU=ON (934183-16-1/BI OR 111119-28-9/BI OR 137076-54-1/BI OR 14265-75-9/BI OR 15750-15-9/BI OR 15757-14-9/BI OR 317809-26-0/BI OR 33507-63-0/BI OR 705283-66-5 /BI OR 901439-51-8/BI OR 901439-89-2/BI OR 901442-07-7/BI OR 901443-47-8/BI OR 91037-65-9/BI OR 934183-14-9/BI OR 934183-15-0/BI OR 934350-78-4/BI OR 934350-82-0/BI OR 934350-86-4/BI OR 934350-87-5/BT OR 10098-91-6/BT OR 110880-55-2/BT OR 110880-57-4/BI OR 111844-19-0/BI OR 112188-16-6/BI OR 115608-61-2/BI OR 118726-52-6/BI OR 128009-23-4/BI OR 135702-31-7/BI OR 137184-55 -5/BI OR 137813-35-5/BI OR 13967-64-1/BI OR 13967-65-2/BI OR 13981-25-4/BI OR 13981-56-1/BI OR 14119-08-5/BI OR 14119-09-6/B I OR 14133-76-7/BI OR 141743-95-5/BI OR 14191-64-1/BI OR 14265-85-1/BI OR 14687-25-3/BI OR 14809-53-1/BI OR 14834-85-6/B I OR 14885-78-0/BI OR 148893-10-1/BI OR 14913-49-6/BI OR 14981-79-4/BI OR 15065-93-7/BI OR 15757-86-5/BI OR 15765-31-8/B I OR 15776-20-2/BI OR 161552-03-0/BI OR 17137-11-0/BI OR 174267-75-5/BI OR 188982-12-9/BI OR 22541-18-0/BI OR 22541-19-1 /BI OR 267410-13-9/BI OR 29022-11-5/BI OR 294-90-6/BI OR 36849-05-5/BI OR 41444-88-6/BI OR 415706-07-9/BI OR 507475-91-4 /BI OR 5292-43-3/BI OR 585531-74-4/BI OR 6066-82-6/BI OR 623575-85-9/BI OR 676544-84-6/BI OR 676544-85-7/BI OR 676553-18 -7/BI OR 676553-19-8/BI OR 7087-68-5/BI OR 713520-27-5/BI OR 728914-72-5/BI OR 728914-74-7/BI OR 7429-91-6/BI OR 7439-91-0/B

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               -0/BI OR 766529-24-2/BI OR 766529-25-3/BI OR 76652
L41
            65 SEA FILE-REGISTRY ABB-ON PLU-ON L40 AND L2
            75 SEA FILE=REGISTRY ABB=ON PLU=ON L40 AND M/ELS
L42
L45
             8 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L41 OR L42) AND L38
L47
               STR
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
Structure attributes must be viewed using STN Express query preparation.
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           142 SEA FILE=REGISTRY ABB=ON PLU=ON L49 AND M/ELS
           203 SEA FILE=REGISTRY ABB=ON PLU=ON L49 NOT L50
L51
L54
            9 SEA FILE=REGISTRY ABB=ON PLU=ON L50 AND Y/ELS
            10 SEA FILE=ZCAPLUS ABB=ON PLU=ON L54
L55
L56
          112 SEA FILE=REGISTRY ABB=ON PLU=ON L50 AND LNTH/PG
           18 SEA FILE=ZCAPLUS ABB=ON PLU=ON L32 OR L37 OR L45 OR L55
L58
       641196 SEA FILE=ZCAPLUS ABB=ON PLU=ON ?TUMOUR?/BI OR ?TUMOR?/BI
L60
L62
        25232 SEA FILE=ZCAPLUS ABB=ON PLU=ON ?SCAFFOLD?/BI
             2 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L51 OR L56) AND L62
L64
            40 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L51 OR L56) AND L60
L65
            8 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L64 OR L65) AND L58
L67
=> s (132 or L37 or L45 or L55 or L67) not L73-L74
L79
           17 (L32 OR L37 OR L45 OR L55 OR L67) NOT (L73 OR L74)
=> d ibib abs hitind hitstr L79 1-17
L79 ANSWER 1 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN
                       2007:1302637 ZCAPLUS Full-text
ACCESSION NUMBER:
DOCUMENT NUMBER:
                        147:522590
TITLE:
                       Preparation of peptides containing the
                        D-Phe-D-Phe-D-Val-D-Leu-D-Lys sequence as imaging
                       agents
INVENTOR(S):
                       Austen, Brian
PATENT ASSIGNEE(S):
                       St. George's Hospital Medical School, UK
SOURCE:
                        PCT Int. Appl., 36pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                       KIND DATE APPLICATION NO. DATE
    PATENT NO.
    WO 2007129077
                        Α2
                             20071115
                                        WO 2007-GB1669
                                                                20070504
     WO 2007129077
                        A3 20080103
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA,
            CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB,
            GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM,
            KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK,
            MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO,
            RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT,
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TZ, UA, UG, US, UZ, VC, VN, ZA. ZM. ZW

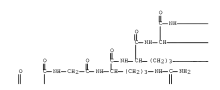
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RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW,
             GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA
PRIORITY APPLN. INFO .:
                                            GB 2006-8960
                                                                A 20060505
     The invention relates to synthetic peptides capable of recognizing and binding
     to B-amyloid and to the use of the peptides in the diagnosis, monitoring and
     therapy of Alzheimer's disease (AD). Peptides containing the sequence D-Phe-
     D-Phe-D-Val-D-Leu-D-Lys (ffylk) and an amine or quanidine substituent are
     claimed for this purpose. Thus, acetyl-rGffvlkr-NH2 and DOTA-rGffvlkGrG-
     pentadiamine (DOTA = 1,4,7,10-tetraazacyclododecane- 1,4,7,10-tetraacetic
     acid) Gd complex were prepared by the solid-phase method and assayed for
     inhibition of \beta-amyloid oligomer formation.
     34-3 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 1, 78
     956489-86-4P 956599-09-0P 956599-10-3P
     RL: DGN (Diagnostic use); SPN (Synthetic preparation); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (preparation of peptides containing D-configurated
phenylalanylphenylalanylvalyl
        leucylleucine sequence as imaging agents)
     956489-89-7P 956489-91-1P 956489-93-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of peptides containing D-configurated
phenylalanylphenylalanylvalyl
        leucylleucine sequence as imaging agents)
     956599-09-0P 956599-10-3P
     RL: DGN (Diagnostic use); SPN (Synthetic preparation); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (preparation of peptides containing D-configurated
phenylalanylphenylalanylvalyl
        leucylleucine sequence as imaging agents)
     956599-09-0 ZCAPLUS
RN
CN
    Gadolinium, [N-[2-[4,7,10-tris[(carboxy-κ0)methyl]-1,4,7,10-
     tetrazacyclododec-1-y1-KN1, KN4, KN7, KN101acetv1-
     κO]-D-arginvlglvcvl-D-phenvlalanvl-D-phenvlalanvl-D-valvl-D-leucvl-D-
     lysylglycyl-D-arginyl-N-(5-aminopentyl)glycinamidato(3-)]- (CA INDEX
    NAME)
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PAGE 1-B

PAGE 2-A

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- RN 956599-10-3 ZCAPLUS
- CN Gadolinium, [N-[2-[4,7,10-tris[(carboxy- κ 0)methyl]-1,4,7,10-tetrazadodec-1-yl- κ N1, κ N4, κ N7, κ N10]acetyl-
 - κ O]-D-arginylglycyl-D-phenylalanyl-D-phenylalanyl-D-valyl-D-leucyl-D-lysylglycyl-D-arginyl-D-



PAGE 1-C

PAGE 2-A

IT 956489-91-1P 956489-93-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of peptides containing D-configurated

phenylalanylphenylalanylvalyl

leucylleucine sequence as imaging agents)

RN 956489-91-1 ZCAPLUS CN Glycinamide, N2-[2-]

Glycinamide, N2-{2-{4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-D-arginylglycyl-D-phenylalanyl-D-phenylalanyl-D-valyl-Dleucyl-D-lysylglycyl-D-arginyl-N-(5-aminopentyl)- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

PAGE 2-B

- RN 956489-93-3 ZCAPLUS
- CN D-Argininamide, N2-[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-D-arginylglycyl-D-phenylalanyl-D-phenylalanyl-D-leucyl-D-lysylglycyl-D-arginyl-D-a

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 2-A

PAGE 2-B

PAGE 3-A

L79 ANSWER 2 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:410019 ZCAPLUS Full-text

DOCUMENT NUMBER: 146:415599

TITLE: Neuropeptide Y analogs for treating and diagnosing Y1

receptor-expressing breast cancer

INVENTOR(S): Srinivasan, Ananth

PATENT ASSIGNEE(S): Schering Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 83pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE:

English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KIND DATE				APPL		DATE						
WO 2007039318				A2 20070412				WO 2	006-	20061005							
WO 2007039318				A3 20070705													
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	KM,	KN,	KΡ,
		KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,
		MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,
		RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	zw							
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		IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	KZ,	MD,	RU,	TJ,	TM,	AP,	EA,	EP,	OA						

PRIORITY APPLN. INFO.: US 2005-723909P P 20051006

The neuropeptide Y(NPY)-receptor-subtype Y1 is expressed differentially from breast tumor cells and is therefore an advantageous target mol. for the mol. imaging of breast cancer. Peptide analogs were synthesized, whose sequence is reduced to the receptor-binding sections of the natural ligand NPY. These Y1 receptor-selective peptide analogs contain unnatural amino acids that increase the receptor affinity and are to ensure the stability of the greatly shortened peptide. New NPY analogs, which are to be used as radioligands, were tested for their binding affinity and selectivity for the Y1 receptor. To this end, in-vitro binding tests with Y1- or Y2 receptor-expressing cell lines were established and optimized. Then, the binding affinities of the NPY analogs were determined. In this case, a peptide (P2489) was identified, whose highest binding affinity was determined with a Ki of 42.8 nmol of Y1 receptor-

expressing SK-M-MC cells and whose selectivity for the Y1 receptor could be detected by the fact that there is no binding to Y2 receptor-expressing MHH-NB-11 cells. As an addnl. NPY analog, peptide fW7 contained the unnatural amino acid β -aminocyclopropanecarboxylic acid on positions 32 and 34, by which the binding to the Y1 receptor was influenced in a pos. manner. A direct coupling of the chelating agent DOTA, which is necessary for the radiometal labeling of the peptides, to the N-terminal end of the peptides resulted in the loss of the binding affinity. By indirect coupling of the DOTA to the peptide fW7 via a spacer, this loss could be reduced, and fW7(DOTA) had a high binding affinity (K1 = 62.8 nmol) similar to P2489.

CC 2-10 (Mammalian Hormones)

IT 13981-56-1D, 18 F, complexes with neuropeptide Y analogs, biological studies 14133-76-7D, 99Tc, metastable, complexes with neuropeptide Y analogs, biological studies 14265-75-9D, complexes with neuropeptide Y analogs, biological studies 15750-15-9D, 1111n, complexes with neuropeptide Y analogs, biological studies 15757-14-9D, complexes with neuropeptide Y analogs, biological studies 15757-14-9D, complexes with neuropeptide Y analogs, biological studies 82785-45-3D, Neuropeptide Y, analogs 705283-66-5D, labeled 934183-14-9D, labeled 934183-15-0D, labeled 934183-15-1D, labeled 934350-87-5D, labeled 934350-87-6D, labeled 934350-87-5D, labeled RL: DGN (Diagnostic use) BIOL (Biological study); USES (Uses) (neuropeptide Y analogs for treating and diagnosing Y receptor-expressing breast cancer)

T 705283-66-5 934183-14-9 934183-15-0 934183-16-1 934350-78-4 934350-87-5 RL: DGN (Diagnostic use), PAC (Pharmacological activity); THU (Therapeutic use), BIOL (Biological study); USES (Uses) (neuropeptide Y analogs for treating and diagnosing Y

receptor-expressing breast cancer)

II 934183-16-10, 177-Lu-DOTA complexes 934350-98-6
RL: DGN (Diagnostic use); PKT (Pharmacokinetics); BIOL (Biological study); USES (Uses)

(neuropeptide Y analogs for treating and diagnosing Y receptor-expressing breast cancer)

II 14133-76-70, 99Tc, metastable, complexes with neuropeptide Y analogs, biological studies 14265-75-9D, complexes with neuropeptide Y analogs, biological studies 15750-15-9D, 111In, complexes with neuropeptide Y analogs, biological studies 15757-14-9D, complexes with neuropeptide Y analogs, biological studies 324183-15-9D, labeled

RL: DGN (Diagnostic use); BIOL (Biological study); USES (Uses) (neuropeptide Y analogs for treating and diagnosing Y

receptor-expressing breast cancer)

RN 14133-76-7 ZCAPLUS

CN Technetium, isotope of mass 99 (CA INDEX NAME)

99Tc

RN 14265-75-9 ZCAPLUS

CN Lutetium, isotope of mass 177 (CA INDEX NAME)

177 Lu

RN 15750-15-9 ZCAPLUS

CN Indium, isotope of mass 111 (CA INDEX NAME)

1111n

RN 15757-14-9 ZCAPLUS

CN Gallium, isotope of mass 68 (CA INDEX NAME)

68Ga

RN 934183-15-0 ZCAPLUS

CN L-Tyrosinamide, N2-[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-trazacyclododec-1-yl]acetyl]-L-arginyl-L-histidyl-L-tyrosyl-L-isoleucyl-L-asparaginyl-L-leucyl-L-isoleucyl-(15,25,35)-2-amino-3-(methoxycarbonyl)cyclopropanecarbonyl-L-arginyl-(15,25,35)-2-amino-3-(methoxycarbonyl)cyclopropanecarbonyl-L-arginyl- (CA INDEX NAME)

PAGE 1-C

IT 934183-15-0

RL: DGN (Diagnostic use); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(neuropeptide Y analogs for treating and diagnosing Y receptor-expressing breast cancer)

- RN 934183-15-0 ZCAPLUS
- CN L-Tyrosinamide, N2-[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-

tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-histidyl-L-tyrosyl-L-isoleucyl-L-asparaginyl-L-leucyl-L-isoleucyl-(15,28,38)-2-amino-3-(methoxycarbonyl)cyclopropanecarbonyl-L-arqinyl-(15,28,38)-2-amino-3-

(methoxycarbonyl)cyclopropanecarbonyl-L-arginyl- (IS, 25, 35)-2-amino-(methoxycarbonyl)cyclopropanecarbonyl-L-arginyl- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-C

IT 934350-88-6

CN

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); BIOL (Biological study); USES (Uses)

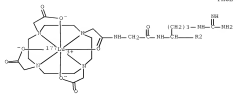
(neuropeptide Y analogs for treating and diagnosing Y receptor-expressing breast cancer)

RN 934350-88-6 ZCAPLUS

Lutetium-177Lu, [N-[2-[4,7,10-tris[(carboxy-K0)methyl]-1,4,7,10-tetraazacyclododec-1-yl-KNl, KN4, KN7, KN10]acetyl-K0]glycyl-L-arginyl-L-histidyl-L-tyrosyl-L-isoleucyl-L-asparaginyl-L-leucyl-L-tisoleucyl-L-threonyl-L-arginyl-2-aminocyclohexanecarbonyl-L-

arginyl-L-tyrosinamidato(3-)]- (CA INDEX NAME)





PAGE 2-A

$$\begin{array}{c|c} & \text{CH}_2-\text{CH}-\text{NH}-\text{C-R} \\ & \text{H}_2\text{N}-\text{C} \end{array}$$

PAGE 3-A

PAGE 3-B

L79 ANSWER 3 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:1274397 ZCAPLUS Full-text

DOCUMENT NUMBER: 147:183765

TITLE: Evaluation of a new biotin-DOTA conjugate for

pretargeted antibody-quided radioimmunotherapy

(PAGRIT)

AUTHOR(S): Urbano, Nicoletta; Papi, Stefano; Ginanneschi, Mauro;

> Santis, Rita; Pace, Silvia; Lindstedt, Ragnar; Ferrari, Liliana; Choi, SunJu; Paganelli, Giovanni;

Chinol, Marco

CORPORATE SOURCE: Division of Nuclear Medicine, European Institute of

Oncology, Milan, 20141, Italy

SOURCE: European Journal of Nuclear Medicine and Molecular

Imaging (2007), 34(1), 68-77 CODEN: EJNMA6; ISSN: 1619-7070

Springer

DOCUMENT TYPE: Journal

PUBLISHER:

LANGUAGE: English AB

Purpose: A novel biotin-DOTA conjugate (r-BHD: reduced biotinamidohexylamine-DOTA) was investigated in order to provide an efficient pretargeted antibodyguided radioimmunotherapy (PAGRIT) application. Preclin. and clin. results are described. Methods: 90Y and 177Lu were used to label r-BHD. The effect of pH and a wide range of specific activities were studied. Radiolabeled r-BHD was tested for affinity towards avidin and for stability in saline or in human serum with and without ascorbic acid. Pharmacokinetic data were collected and organ biodistribution evaluated in a tumor-bearing pretargeted animal model. A pilot study was performed in a metastatic melanoma patient and dosimetry was estimated Results: High radiochem. purity (>99%) was routinely achieved with 90Y or 177Lu in sodium acetate buffer (1.0 M, pH 5.0) at a specific activity of 2.6 MBq/nmol. Both 90Y- and 177Lu-r-BHD were also prepared at higher specific activities. Radiolabeled r-BHD was stable up to 96 h in human serum and saline with the addition of ascorbic acid. The structural modifications proposed for the r-BHD stabilized it against enzymic degradation while retaining high binding affinity for avidin. Renal clearance appeared to be the main route of excretion in animals, and high tumor uptake was observed in the pretargeted animals. The patient study showed a total body clearance of .apprx.85% in 24 h, with a kidney absorbed dose of 1.5 mGy/MBq. Tumor uptake was rapid and the calculated dose to a 10-mm tumor lesion was .apprx.12 mGy/MBq. Conclusion: These results indicate that the new biotin-DOTA conjugate may be a suitable candidate for pretargeting trials. 8-9 (Radiation Biochemistry)

58-85-5D, DOTA conjugates, Lu-177 complexes 14265-75-9D, 177Lu, complexes with DOTA-biotin, biological studies 60239-18-1D, DOTA, biotin

conjugates, Lu-177 complexes 586962-90-5

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(evaluation of new biotin-DOTA conjugate for pretargeted antibody-guided radioimmunotherapy)

IT 586963-90-5

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(evaluation of new biotin-DOTA conjugate for pretargeted antibody-guided radioimmunotherapy)

RN 586962-90-5 ZCAPLUS

CN Yttrium=90Y, [10-[2-[[6-[[5-(hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl)pentyl]amino]hexyl]amino]-2-(oxo-k0]ethyl]-1,4,7,10-tetrazacyclododecane-1,4,7-triacetato(3-)-kN1,kN4,kN7,.kappa.N10,kO1,ko4,kO7]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 45 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 4 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:734439 ZCAPLUS Full-text

DOCUMENT NUMBER: 145:195598

TITLE: Compounds having RD targeting motifs

INVENTOR(S): Achilefu, Samuel

PATENT ASSIGNEE(S): Washington University, USA SOURCE: PCT Int. Appl., 105 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	KIN	D	DATE		1	APPLICATION NO.						DATE				
					_									-		
WO 2006	0789	14		A1		2006	0727	1	WO 2	006-	US20	56		2	0060	120
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	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,
	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
	MZ,	NA,	NG,	NΙ,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
	SG,	SK,	SL,	SM,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,
	VN,	YU,	ZA,	ZM,	zw											

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO.: US 2005-645816P P 20050123

OTHER SOURCE(S): MARPAT 145:195598

- The present invention provides compds. that have motifs that target the compds to cells that express integrins. In particular, the compds have peptides with one or more RD motifs conjugated to an agent selected from an imaging agent and a targeting agent. The compds may be used to detect, monitor and treat a variety of disorders mediated by integrins.
- 63-5 (Pharmaceuticals)
- 111119-28-9D, DTPA 91037-65-9D, conjugates with cypate and glucosamine 111844-19-0D, conjugates with cypate and octreotate conjugates 317809-26-0, Cypate 317809-26-0D, Cypate, conjugates with peptides 901439-51-8D, DTPA conjugates 901442-07-7D, conjugates with cypate and glucosamine 901442-72-6 901442-80-6 901442-87-3 901442-94-2 901443-01-4 901443-47-8 901443-47-8D, conjugates with peptides 901443-61-6 901443-68-3 901443-74-1 901443-82-1 901443-89-8 901443-96-7 901444-04-0 901444-12-0 901444-20-0D, DTPA conjugates 901444-27-7D, DTPA conjugates 901444-34-6D, DTPA conjugates 901444-41-5D, DTPA conjugates 901444-63-1 901444-71-1 901444-79-9 901444-86-8 901445-34-9 901445-41-8 901445-48-5 RL: DGN (Diagnostic use); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(diagnostic and therapeutic peptide conjugates targeted to integrin-pos. cells)

IT 901444-63-1 901444-71-1

RL: DGN (Diagnostic use); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(diagnostic and therapeutic peptide conjugates targeted to

integrin-pos. cells) RN 901444-63-1 ZCAPLUS

CN L-Lysine, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginylglycyl-L-α-aspartyl-L-seryl-L-prolyl- (9CI) (CA INDEX NAME)

RN 901444-71-1 ZCAPLUS

CN L-Lysinamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginylglycyl-L-a-aspartyl-L-seryl-L-prolyl-M6-[3-[2-[7-[3-(2-carboxyethyl)-1,3-dihydro-1,1-dimethyl-2H-benz[e]indol-2-ylidene]-1,3,5-heptatrienyl]-1,1-dimethyl-1H-benz[e]indolio]-1-oxopropyl]-, bromide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

PAGE 1-A



PAGE 2-B

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 5 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:625378 ZCAPLUS Full-text

DOCUMENT NUMBER: 145:243952

TITLE: Magnetic resonance imaging of tumor cells by

targeting the amino acid transport system AUTHOR(S): Lattuada, Luciano; Demattio, Silvia; Vincenzi,

Veronica; Cabella, Claudia; Visigalli, Massimo; Aime, Silvio; Crich, Simonetta Geninatti; Gianolio, Eliana

CORPORATE SOURCE: CRM Chemistry, Bracco Imaging SpA, Milan, 20134, Italy SOURCE: Bioorganic & Medicinal Chemistry Letters (2006),

16(15), 4111-4114

CODEN: BMCLE8: ISSN: 0960-894X

PUBLISHER: Elsevier B.V. DOCUMENT TYPE: Journal

LANGUAGE: English OTHER SOURCE(S): CASREACT 145:243952

An early diagnosis of cancer is crucial in the battle against this disease and the in vivo visualization of tumor's at cellular level is still the most challenging goal. In order to target tumor cells, we took into account their increased metabolism and amino acid nutrients or pseudo-nutrients, which are actively transported through the cell membrane, have been chosen as vectors for new MRI contrast agents. For this reason new gadolinium complexes conjugated to agmatine, arginine, and glutamine have been synthesized and studied.

8-9 (Radiation Biochemistry)

MRI tumer aminoacid transport prepn gadolinium complex conjugate; agmatine arginine glutamine conjugate gadolinium MRI contrast agent

Neoplasm (MRI of tumor by targeting amino acid transport: preparation of

gadolinium complexes conjugated to agmatine, arginine, and glutamine)

Imaging agents

(NMR contrast; MRI of tomer by targeting amino acid

transport: preparation of gadolinium complexes conjugated to agmatine, arginine, and glutamine)

Imaging

(NMR; MRI of tumor by targeting amino acid transport: preparation

of gadolinium complexes conjugated to agmatine, arginine, and $\operatorname{qlutamine}$)

IT Imaging

(tumor; MRI of tumor by targeting amino acid

transport: preparation of gadolinium complexes conjugated to agmatine, arginine, and glutamine)

IT 906078-01-1P 906078-02-2P 906078-03-3P 906078-04-4P 906078-05-5P 906078-06-6P

906078-07-7P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(MRI of tumor by targeting amino acid transport: preparation of gadolinium complexes conjugated to agmatine, arginine, and glutamine)

T 79-04-9, Chloroacetylchloride 6066-82-6, N-Hydroxysuccinimide 41444-88-6 115608-61-2 128009-23-4 174267-75-5 585531-74-4 805233-27-6

RL: RCT (Reactant); RACT (Reactant or reagent)

(MRI of tumor by targeting amino acid transport: preparation of

gadolinium complexes conjugated to agmatine, arginine, and glutamine) IT 905985-29-7P 905985-30-0P 905985-31-1P 905985-32-2P 905985-34-4P 905985-35-5P 905985-36-6P 905985-37-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(MRI of tumor by targeting amino acid transport: preparation of gadolinium complexes conjugated to agmatine, arginine, and glutamine) 906078-01-1P 906078-02-2P 906078-03-3P

906078-04-4P 906078-05-5P 906078-06-6P

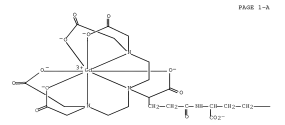
906078-07-7P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(MRI of tumor by targeting amino acid transport: preparation of gadolinium complexes conjugated to agmatine, arginine, and glutamine)

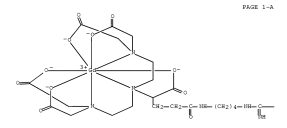
RN 906078-01-1 ZCAPLUS

CN Gadolinate(3-), [N,N-bis[2-[bis[(carboxy-x0)methyl]aminokN]ethyl]-L-y-glutamyl-xN,XOI-L-glutaminato(6-)]-, trisodium (9CI) (CA INDEX NAME)



●3 Na+

- RN 906078-02-2 ZCAPLUS
- CN Gadolinate(2-), [1-amino-12-[2-[bis[(carboxy-k0)methyl]amino-kN]ethyl]-11-(carboxy-k0)-15-[(carboxy-k0)methyl]-1-imino-8-oxo-2,7,12,15-tetraazaheptadecan-17-oato(5-)-kN12,kN15,kO17]-, disodium (9CI) (CA INDEX NAME)

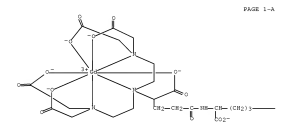


●2 Na+

--- NH2

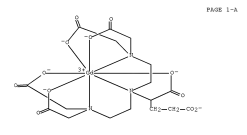
RN 906078-03-3 ZCAPLUS

CN Gadolinate(3-), [N,N-bis[2-[bis[(carboxy-k0)methyl]aminokN]ethyl]-L-y-glutamyl-kN, kOl-L-argininato(6-)]-, trisodium (9CI) (CA INDEX NAME)



RN 906078-04-4 ZCAPLUS

CN Gadolinate(3-), [N,N-bis[2-[bis[(carboxy-K0)methyl]amino-KN]ethyl]-L-glutamato(6-)-KN2,KOl]-, trisodium (9CI) (CA INDEX NAME)



PAGE 2-A

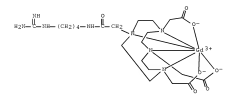
■3 Na+

- RN 906078-05-5 ZCAPLUS
- CN Gadolinate(1-), [10-[2-[(4-amino-1-carboxy-4-oxobuty1) amino]-2-oxoethy1]1,4,7,10-tetraazazoylododecane-1,4,7-triacetato(4-)xN1,xN4,xN7,xN10,x01,x04,x07]-,

hydrogen (9CI) (CA INDEX NAME)

RN 906078-06-6 ZCAPLUS

CN Gadolinium, [10-[2-[[4-[(aminoiminomethyl)amino]butyl]amino]-2-oxoethyl]1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(7-)KN1,KN4,KN7,KN10,KO1,KO4,KO7](9CI) (CA INDEX NAME)



RN 906078-07-7 ZCAPLUS

CN Gadolinate(1-), [10-[2-[[4-[(aminoiminomethyl)amino]-1-carboxybutyl]amino]-2-oxoethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(4-)kN1,kN4,kN7,kN10,kO1,kO4,kO7]-,
hydrogen (9C1) (CA INDEX NAME)

905985-35-5P 905985-37-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(MRI of temor by targeting amino acid transport: preparation of gadolinium complexes conjugated to agmatine, arginine, and glutamine)

905985-35-5 ZCAPLUS RN CN

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[4-[(aminoiminomethyl)amino]butyl]amino]-2-oxoethyl]- (CA INDEX NAME)

905985-37-7 ZCAPLUS

ĊN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[(1S)-4-[(aminoiminomethyl)amino]-1-carboxybutyl]amino]-2-oxoethyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMA.

L79 ANSWER 6 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2005:1289862 ZCAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 144:31701

TITLE: Preparation of metal complexes of trimeric

DOTA-macrocyclic substituted aminoisophthalate

trihalophenyl derivatives

INVENTOR(S): Harto, Juan R.; Martin, Jose L.; Platzek, Johannes;

Schirmer, Heiko; Weinmann, Hanns-Joachim; Carretero,

Jose
PATENT ASSIGNEE(S): Schering Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PA	TENT	NO.			KIND DATE					APPL								
WO	2005	1159	97		A1 20051208			WO 2005-EP4493						20050422				
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		CN,	CO,	CR,	CU,	CZ,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	GE,	
		GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KP,	KR,	KZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	ΝI,	
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	
		SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KΕ,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
		AZ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,	
		RO,	SE,	SI,	SK,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	
		MR,	ΝE,	SN,	TD,	TG												
DE	1020	0402	6103		A1		2005	1222		DE 2	004-	1020	0402	6103	2	0040	525	
EP	1748	992			A1		2007	0207		EP 2	005-	7410	25		2	0050	422	
	R:							DE,								HU,	ΙE,	
				LI,				NL,										
	2008							0110										
US	2006	1209	65		A1		2006	0608										
PRIORIT	Y APP	LN.	INFO	.:						DE 2	004-	1020	0402	6103.	A 2	0040	525	
										US 2	004-	5754	17P		P 2	0040	601	
										WO 2	005-	EP44	93		W 2	0050	422	
										US 2	005-	1356	56		A1 2	0050	524	
OTHER S	OURCE	(S):			MARI	PAT	144:	3170	1									

OTHER SOURCE(S): MARPAT 144:31701

AB The preparation is described for metal complexes of trihalobenzene functionalized with three DOTA-like chelating groups (I), where X = bromo or iodo. These complexes are suitable as contrast agents. Thus, the ligand I (X = iodo) was prepared in a multistep procedure and was used to prepare Gd, Dy, Yb and Y complexes.

Ι

- IC ICM C07D257-02
 - ICS A61K049-04; A61K051-04; A61K049-08; C07K005-023; C07K005-02
- CC 78-7 (Inorganic Chemicals and Reactions) Section cross-reference(s): 8, 28
- IT 7429-91-6P, Dysprosium, preparation 7439-89-6P, Iron, preparation 7440-56-5P, Manganese, preparation 7440-54-2P, Gadolinium, preparation 870475-42-6P 870475-43-7P

870475-44-8P 870475-45-9P 870475-48-2DP, metal complexes RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological

study); PREP (Preparation); USES (Uses)

(preparation of metal complexes with trihalobenzene functionalized with three DOTA-like chelating groups for use as contrast agents)

IT 870475-45-9P

RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of metal complexes with trihalobenzene functionalized with three DOTA-like chelating groups for use as contrast agents)

RN 870475-45-9 ZCAPLUS

CN Yttrium, [μ3-[[10,10'-[[2,4,6-triiodo-5-[methyl[[[1-(οxο-κ0)-2-[4,7,10-tris[(carboxy-κ0)methyl]-1,4,7,10-terraazacyclododec-1-yl-κN1,κN4,N7,N10]propyl]amino]acetyl]amino]-1,3-

) | | tri- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 2-A

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REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 7 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2005:1220695 ZCAPLUS Full-text

DOCUMENT NUMBER: 143:471966

TITLE: Macrocycle-substituted trimer halogen-benzene

derivatives

INVENTOR(S): Harto, Juan R.; Martin, Jose L.; Platzek, Johannes; Schirmer, Heiko; Weinmann, Hanns-Joachim; Carretero,

Jose

PATENT ASSIGNEE(S): Schering Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 49 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

WO 2005108379 Al 20051117 WO 2005-EP4319 2005. W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BB, BR, BR, BW, BY, BZ, CC, CR, CO, CR, CU, CZ, DK, DM, DZ, EC, EE, EG, ES, FI, GB, CG, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, F, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MM, MW, MX, MZ, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZR: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZA, BY, KG, KZ, MD, RU, TJ, TR, AT, BE, BG, CH, CY, CZ, I				
CN, CO, CR, CU, CZ, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GH, GM, HR, HJ, ID, IL, IN, IS, JP, KE, KG, KM, KF, KR, F, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MM, MX, MZ, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZRW, BW, GH, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, Z	419			
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LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, N NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, S SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, VY, RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, 2	GE,			
NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, S SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, Z RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, Z	LC,			
SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, Z RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, Z	NI,			
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	ZM, ZW			
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	DK,			
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RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, G	ML,			
MR, NE, SN, TD, TG				
DE 102004023093 B3 20060302 DE 2004-102004023093 200	505			
EP 1742926 A1 20070117 EP 2005-742880 200	419			
EP 1742926 B1 20070808				
R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, F	IE,			
IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR				
AT 369347 T 20070815 AT 2005-742880 200	419			
JP 2007536295 T 20071213 JP 2007-511925 200	419			
ES 2289711 T3 20080201 ES 2005-742880 200				

A1 20060713 US 2006154989 HS 2005-272008 20051114 PRIORITY APPLN. INFO.: DE 2004-102004023093A 20040505 US 2004-574713P P 20040527 WO 2005-EP4319 W 20050419 US 2005-122248 A1 20050505

OTHER SOURCE(S): MARPAT 143:471966

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to rare earth, Fe and Mn complexes of I (X = Br or I' Al AB = CONR1(CH2)nNR2(COCHZ1NH)mCOCHZ2K, A2 = NR1COCHZ2K (R1 and R1 = H, C1-2 alky1 group of monohydroxy C1-2 alkyl group; Z1 and Z2 = H or Me; n = 2-4; m = 0-1; K = 1,4,7,11-tetraazacyclotetradecane-1,4,7-triacetic acid group)) and said complexes are suitable as contrast agents. For example, II (H3L) was prepared in a multi step process starting from 2,4,6-triiodo-5-(methylamino) isophthaloyl dichloride and ethylenediamine, with subsequent reaction with 2-bromopropanoyl bromide, 1,4,7-tris(benzylcarbonyl)-1,4,7,11tetraazacyclotetradecane with deprotection and reaction with chloroacetic acid. GdL in 58 % yield was prepared from II and Gd203.

TC: ICM C07D257-02

ICS A61K051-04; A61K049-08

78-7 (Inorganic Chemicals and Reactions) Section cross-reference(s): 9, 28, 77

ΙT 7429-91-6DP, Dysprosium, complexes with tetraazacvclotetradecanetriacetic acid, isophthalic acid amide derivs. 7439-89-6DP, Iron, complexes with tetraazacvclotetradecanetriacetic acid, isophthalic acid amide derivs. 7439-96-5DP, Manganese, complexes with tetraazacyclotetradecanetriacetic acid, isophthalic acid amide derivs. 7440-53-1DP, Europium, complexes with tetraazacyclotetradecanetriacetic acid, isophthalic acid amide 7440-54-2DP, Gadolinium, complexes with tetraazacyclotetradecanetriacetic acid, isophthalic acid amide derivs. 869339-24-2P 869339-25-3P 869339-26-4P 869339-28-6P 869339-51-5DP, isophthalic acid amide derivs., transition metal complexes RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation as contrast agents) TT 869339-26-4P

RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation as contrast agents)

RN 869339-26-4 ZCAPLUS

CN Yttrium, $[\mu 3 - [[10, 10' - [[2, 4, 6 - triiodo - 5 - [methyl [1 - (oxo - <math>\kappa 0) - 2 -$ [4,7,10-tris[(carboxy-KO)methyl]-1,4,7,10-tetraazacyclododec-1-yl-KN1, KN4, KN7, KN10|propvl|amino|-1,3-

phenylene|bis(carbonylimino-2,1-ethanediylimino(1-methyl-2-(οxο-κ0)-2,1-ethanediyl]]]bis[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-KN1, KN4, KN7, KN10, KO1, KO4, KO711(9-

) | | tri- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 2-A

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PAGE 2-B

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 8 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2004:799481 ZCAPLUS Full-text

DOCUMENT NUMBER: 141:320007

TITLE: Radiopharmaceuticals for cancer diagnosis and

treatment

INVENTOR(S): Merlo, Adrian; Maecke, Helmut; Reubi, Jean-Claude;

Good, Stephan

PATENT ASSIGNEE(S): Kantonsspital Basel, Switz.; Universitaet Bern

SOURCE: PCT Int. Appl., 37 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.						_	DATE APPLICATION NO.												
WO	2004	0827	22				2004	0930								0040			
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH.		
		CN,	co,	CR,	CU,	CZ,	DE,	DK.	DM,	DZ,	EC,	EE,	EG,	ES,	FI.	GB,	GD.		
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		LK,	LR,	LS.	LT.	LU,	LV.	MA,	MD,	MG,	MK,	MN.	MW,	MX,	MZ,	NA.	NI		
		NO.	NZ,	OM,	PG.	PH,	PL,	PT.	RO,	RU,	sc.	SD,	SE,	SG,	SK,	SL,	SY		
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EΡ					A1		2004	0922		EP 2	003-		20030319						
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ΑIJ	2004														318				
									CA 2004-2519315										
										EP 2004-721547									
							ES,												
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JP 2007527366 T 20070927 JP 2006-505475 20040318
US 2007053837 A1 20070308 US 2005-549665 20050919
PRIORITY APPLN. INFO:: EP 2003-6061 A 20030319
WO 2004-EP50329 W 20040318

OTHER SOURCE(S): MARPAT 141:320007

- AB The invention relates to radiopharmaceutical carriers consisting of a radiolabeled substance P analog conjugated to a chelating agent such as DOTAGA, DOTAGA or DOTA, which are useful for targeting and treatment of brain tumors, especially gliomas.
- IC ICM A61K051-00
- CC 63-5 (Pharmaceuticals)
- Section cross-reference(s): 8
 - 10098-91-6D, Yttrium 90, substance P-conjugated complexes, biological studies 13967-64-15, Dysprosium 165, substance P-conjugated complexes, biological studies 13967-65-2D, Holmium 166, substance P-conjugated complexes, biological studies 13981-25-4D, Copper 64, substance P-conjugated complexes, biological studies 14119-08-5D, Gallium 66, substance P-conjugated complexes, biological studies 14119-03-6D, Gallium 67, substance P-conjugated complexes, biological studies 14191-64-1D, Praseodymium 142, substance P-conjugated complexes, biological studies 14265-75-9D, Lutetium 177, substance P-conjugated complexes, biological studies 14265-85-1D, Actinium 225, substance P-conjugated complexes, biological studies 14687-25-3D, Lead 203, substance P-conjugated complexes, biological studies 14809-53-1D, Yttrium 86, substance P-conjugated complexes, biological studies 14834-85-6D. Dysprosium 162, substance P-conjugated complexes, biological studies 14885-78-0D, Indium 113, substance P-conjugated complexes, biological studies 14913-49-6D, Bismuth 212, substance P-conjugated complexes, biological studies 14931-79-4D, Praseodymium 143, substance P-conjugated complexes, biological studies 15065-93-7D, Terbium 149, substance P-conjugated complexes, biological studies 15750-15-9D, Indium 111, substance P-conjugated complexes, biological studies 15757-14-9D, Gallium 68, substance P-conjugated complexes, biological studies 15757-86-5D, Copper 67, substance P-conjugated complexes, biological studies 15765-31-8D, Promethium 149, substance P-conjugated complexes, biological studies 15776-20-2D, Bismuth 213, substance P-conjugated complexes, biological studies 33507-63-0D, Substance P, conjugates of radionuclide complexes 36849-05-5D, Dysprosium 167, substance P-conjugated complexes, biological studies 77128-75-7D, conjugates of radionuclide complexes 110880-55-2D, conjugates of radionuclide complexes 110880-57-4D, conjugates of radionuclide complexes 766529-14-0D, conjugates of radionuclide complexes 766529-15-1D, conjugates of radionuclide complexes 766529-16-2D, conjugates of radionuclide complexes 766529-18-4D, conjugates of radionuclide complexes 766529-19-5D, conjugates of radionuclide complexes 766529-20-8D, conjugates of radionuclide complexes 766529-22-0D, conjugates of radionuclide complexes 766529-24-2D, conjugates of radionuclide complexes 766529-25-3D, conjugates of radionuclide complexes 766529-28-6D, conjugates of radionuclide complexes 766529-29-7D, conjugates of radionuclide complexes RL: DGN (Diagnostic use); THU (Therapeutic use); BIOL (Biological study);
 - KL: DEN (Diagnostic use); THO (Inerapeutic use); BIOL (Biological study).
 USES (Uses)

 (radiolabeled substance P conjugates for cancer diagnosis and
 - treatment)
- IT 767340-53-4P
 - RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

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     766529-36-6P 766529-37-7P 766529-38-8P
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    biological studies 15765-31-80, Promethium 149, substance
     P-conjugated complexes, biological studies 15776-20-2D, Bismuth
     213, substance P-conjugated complexes, biological studies
     36849-05-50, Dysprosium 167, substance P-conjugated complexes,
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     RL: DGN (Diagnostic use); THU (Therapeutic use); BIOL (Biological study);
     USES (Uses)
       (radiolabeled substance P conjugates for cancer diagnosis and
       treatment)
RN
     10098-91-6 ZCAPLUS
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Yttrium, isotope of mass 90 (CA INDEX NAME)

90Y

CN

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RN
   13967-64-1 ZCAPLUS
CN Dysprosium, isotope of mass 165 (CA INDEX NAME)
165Dy
RN 13967-65-2 ZCAPLUS
CN Holmium, isotope of mass 166 (CA INDEX NAME)
166Ho
RN 13981-25-4 ZCAPLUS
CN
    Copper, isotope of mass 64 (CA INDEX NAME)
64cu
   14119-08-5 ZCAPLUS
CN Gallium, isotope of mass 66 (CA INDEX NAME)
66Ga
RN
   14119-09-6 ZCAPLUS
CN Gallium, isotope of mass 67 (CA INDEX NAME)
67ga
RN
   14191-64-1 ZCAPLUS
CN Praseodymium, isotope of mass 142 (CA INDEX NAME)
142pr
RN 14265-75-9 ZCAPLUS
CN Lutetium, isotope of mass 177 (CA INDEX NAME)
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10/573938
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 $177 \mathrm{Lu}$

RN 14265-85-1 ZCAPLUS

CN Actinium, isotope of mass 225 (CA INDEX NAME)

225Ac

RN 14687-25-3 ZCAPLUS

CN Lead, isotope of mass 203 (CA INDEX NAME)

203Pb

RN 14809-53-1 ZCAPLUS

CN Yttrium, isotope of mass 86 (CA INDEX NAME)

86Y

RN 14834-85-6 ZCAPLUS

CN Dysprosium, isotope of mass 162 (CA INDEX NAME)

162Dy

RN 14885-78-0 ZCAPLUS

CN Indium, isotope of mass 113 (CA INDEX NAME)

113_I

RN 14913-49-6 ZCAPLUS

CN Bismuth, isotope of mass 212 (CA INDEX NAME)

212Bi

- RN 14981-79-4 ZCAPLUS
- CN Praseodymium, isotope of mass 143 (CA INDEX NAME)

143pr

- RN 15065-93-7 ZCAPLUS
- CN Terbium, isotope of mass 149 (CA INDEX NAME)

149 Tb

- RN 15750-15-9 ZCAPLUS
- CN Indium, isotope of mass 111 (CA INDEX NAME)

1111n

- RN 15757-14-9 ZCAPLUS
- CN Gallium, isotope of mass 68 (CA INDEX NAME)

68_{Ga}

- RN 15757-86-5 ZCAPLUS
- CN Copper, isotope of mass 67 (CA INDEX NAME)

67cu

- RN 15765-31-8 ZCAPLUS
- CN Promethium, isotope of mass 149 (CA INDEX NAME)

149 pm

- RN 15776-20-2 ZCAPLUS
- CN Bismuth, isotope of mass 213 (CA INDEX NAME)

213Bi

RN 36849-05-5 ZCAPLUS

CN Dysprosium, isotope of mass 167 (CA INDEX NAME)

167Dv

IT 767340-53-4P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(radiolabeled substance P conjugates for cancer diagnosis and treatment)

RN 767340-53-4 ZCAPLUS

CN Indate(1-)-111In, [N2-[4-(carboxy-KO)-1-oxo-4-[4,7,10-tris[(carboxy-KO)methyl]-1,4,7,10-tetraazacyclododec-1-ylKN1,KN4,KN7,KN10]butyl]-L-arginyl-L-prolyl-L-lysylL-prolyl-L-glutaminyl-L-glutaminyl-L-phenylalanyl-L-phenylalanyl-lleucyl-L-methioninamidato(4-)]- (9C1) (CA INDEX NAME)

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IT 766529-31-1P 766529-32-2P 766529-33-3P 766529-34-4P 766529-35-5P 766529-36-6P

766529-37-7P 766529-38-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(radiolabeled substance P conjugates for cancer diagnosis and

treatment)

- RN 766529-31-1 ZCAPLUS
- CN Substance P, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-11-[(2S)-2-amino-4-(methylsulfonyl)butanamide]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

- RN 766529-32-2 ZCAPLUS
- CN L-Methioninamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-phenylalanyl-L-phenylalanyl-N-methylglycyl-L-leucyl- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

- RN 766529-33-3 ZCAPLUS
- CN L-Methioninamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-phenylalanyl-3-(2-thienyl)-L-alanylglycyl-L-leucyl- (9CI) (CA INDEX NAME)

RN 766529-34-4 ZCAPLUS

CN L-Methioninamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-glutaminyl-3-(2-thienyl)-L-alanyl-L-phenylalanylglycyl-L-leucyl- (9CI) (CA INDEX NAME)

RN 766529-35-5 ZCAPLUS

CN Substance P, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-9-(N-methylglycine)-11-[(2S)-2-amino-4-(methylsulfonyl)butanamide]- (9CI) (CA INDEX NAME)

- RN 766529-36-6 ZCAPLUS
- CN Butanamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-phenylalanyl-3-(2-thienyl)-L-alanylglycyl-L-leucyl-2-amino-4-(methylsulfonyl)-, (25)- (9CI) (CA INDEX NAME)

- RN 766529-37-7 ZCAPLUS
- CN L-Methioninamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yllacetyl]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-phenylalanyl-3-(2-thienyl)-L-alanyl-N-methylglycyl-L-leucyl- (9CI) (CA INDEX NAME)

RN 766529-38-8 ZCAPLUS

CN L-Methioninamide, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yllacetyl]-L-aginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-3-(2-thienyl)-L-alanyl-3-(2-thienyl)-L-alanylglycyl-L-leucyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 767340-54-5P 767340-55-6P 767340-56-7P 767340-57-8P 767340-58-9P 767340-53-0P

767340-57-8P 767340-58-9P 767340-59-0P 767340-60-3P 767340-61-4P 767340-62-5P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

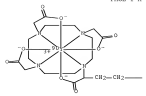
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RN 767340-54-5 ZCAPLUS

CN Yttrate(1-)-90Y, [N2-[4-(carboxy- κ 0)-1-oxo-4-[4,7,10-tris[(carboxy- κ 0)methyl]-1,4,7,10-tetraazacyclododec-1-yl-

KN1, KN4, KN7, KN10]butyl]-L-arginyl-L-prolyl-L-lysyl-

L-prolyl-L-glutaminyl-L-glutaminyl-L-phenylalanyl-L-phenylalanylglycyl-L-leucyl-L-methioninamidato(4-)]- (9CI) (CA INDEX NAME)



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RN 767340-55-6 ZCAPLUS

CN Indium-111In, [N2-[[4,7,10-tris[(carboxy-K0)methyl]-1,4,7,10-tetraazacyclododec-1-yl-KN1,KN4,KN7,KN10]acetyl-K0]-11-[[2S)-2-amino-4-(methylsulfonyl)butanamide]substance P-ato(3-)]-(9CI) (CA INDEX NAME)

PAGE 1-A

- RN 767340-56-7 ZCAPLUS
- CN Indium-111In, [N2-[[4,7,10-tris[(carboxy- κ 0)methy1]-1,4,7,10-tetraazacyclododec-1-y1- κ N1, κ N4, κ N7, κ N10]acety1
 - k0]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-glutaminyl-Lphenylalanyl-L-phenylalanyl-N-methylglycyl-L-leucyl-L-methioninamidato(3-)]- (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 767340-57-8 ZCAPLUS

CN Indium-111In, [N2-[[4,7,10-tris]((carboxy-K0)methyl]-1,4,7,10tetraazacyclododec-l-yl-kNl,kN4,kN7,kNl0]acetylK0]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-Lphenylalanyl-3-(2-thienyl)-L-alanylglycyl-L-leucyl-L-methioninamidato(3-)](9CI) (CA INDEX NAME)

PAGE 1-A

RN 767340-58-9 ZCAPLUS

CN Indium-11lIn, [N2-[[4,7,10-tris[(carboxy-K0)methyl]-1,4,7,10tetrazacyclododec-1-yl-kNl,kN4,kN7,kNl0]acetylk0]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-glutaminyl-3(2-thienyl)-L-alanyl-L-phenylalanylglycyl-L-leucyl-L-methioninamidato(3-)](9CI) (CA INDEX NAME)

PAGE 2-A

RN 767340-59-0 ZCAPLUS

CN Indium-lllIn, [N2-[[4,7,10-tris[(carboxy-KO)methyl]-1,4,7,10-tetraazacyclododec-l-yl-KNl,KNl,KNl,KNl,KNlO]acetyl-KO]-9-(N-methylglycine)-11-[(2S)-2-amino-4-(methylsulfonyl)butanamide]substance P-ato(3-)]- (9CI) (CA INDEX NAME)

PAGE 1-A

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RN 767340-60-3 ZCAPLUS

CN Indium-111In, [N2-[[4,7,10-tris[(carboxy-K0)methyl]-1,4,7,10-tetraazacyclododec-1-yl-KN1,KN4,KN7,KN10]acetyl-

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PAGE 1-A

PAGE 2-A

- RN 767340-61-4 ZCAPLUS
- CN Indium-11lIn, [N2-[[4,7,10-tris[(carboxy-K0)methyl]-1,4,7,10tetrazacyclododec-1-yl-kNl,kNl,kNl,kNl0]acetylk0]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-glutaminyl-Lphenylalanyl-3-(2-thienyl)-L-alanyl-N-methylglycyl-L-leucyl-c2s)-2-amino-4(methylsulfonyl)butanamidato(3-)]-(9C1) (CA INDEX NAME)

PAGE 2-A

RN 767340-62-5 ZCAPLUS

CN Indium-111In, [N2-[[4,7,10-tris](carboxy-k0)methyl]-1,4,7,10tetraazacyclododec-1-yl-kNl,kNl,kNl,kNl) (acetylk0]-L-arginyl-L-prolyl-L-lysyl-L-prolyl-L-glutaminyl-L-glutaminyl-3-(2-thienyl)-L-alanyl-3-(2-thienyl)-L-alanylglycyl-L-leucyl-Lmethioninamidato(3-)]- (9CI) (CA INDEX NAME)

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PAGE 1-B

PAGE 2-A

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PAGE 2-B

L79 ANSWER 9 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2004:718526 ZCAPLUS Full-text DOCUMENT NUMBER: 141:243575

TITLE:

Preparation of 1,3,5-trihalo-2,4,6-

benzenetricarboxamide N,N,N-tristetraazacyclododecane metal complexes and related compounds as contrast

media.

INVENTOR(S): Platzek, Johannes; Weinmann, Hanns-Joachim; Schirmer,

Heiko; Martin, Jose Luis; Harto, Juan R.; Riefke,

Bioern

PATENT ASSIGNEE(S): Schering Aktiengesellschaft, Germany

PCT Int. Appl., 103 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: German FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

SOURCE:

PATENT NO.				KIND DATE			APPLICATION NO.						DATE			
WO 2004074267				A1 20040902			WO 2003-EP14149						20031212			
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Hal Al Q1 = NH

- AB Title compds. [I; Hal = Br, iodo; Al = CONR1(CH2)nNR2(COCHZ1NH)mCOCHZ2K, CONR1 (CH2) p (CONR2CH2) mCH (OH) CH2K, CH2O (CH2) pCH (OH) CH2K, CH20 (CH2) nNR1 (COCHZ1NH) mCOCHZ2K, CH2NR1CO (CHZ1NHCO) mCHZ2K; A2 = A1, NR1CO(NR1)m(CH2)pNR2(COCHZ1NH)mCOCHZ1K; R1, R2 = H, alkyl, hydroxyalkyl; Z1, Z2 = H, Me; n = 2-4; m = 0, 1; p = 1-4; K = Q1; X = H, metal ion of element nos, 20-29, 39, 42, 44, 57-83; ≥ 2 X = metal ions], were prepared Thus, 2,4,6triiodo-1,3,5-benzenetricarbonyl trichloride in THF was added to ethylenediamine in THF over 1 h followed by stirring for 14 h to give 70% 2,4,6-triiodo-1,3,5-benzenetricarboxvlic acid tris(2-aminoethvl)amide. This was added to a mixture prepared from the Gd complex of 10-[4-carboxy-1-methyl-2-oxo-3-azabutyl]-1,4,7,10- tetraazacyclododecane-1,4,7-triacetic acid, DCC, and N-hydroxysuccinimide in Me2SO to give 73% 2,4,6-triiodo-1,3,5benzenetricarboxylic acid N,N,N-tris-[3,6-diaza-4,7-dioxo-8-methyloctan-1,8div1-[10-[1,4,7-tris(carboxymethy1)-1,4,7,10-tetraazacyclododecane, Gd complex]]]amide. The latter was used for CT imaging of rat blood vessels and kidnevs.
 - ICM C07D257-02
 - ICS A61K049-04; A61K049-06; A61K051-04; A61K049-08
- CC 28-23 (Heterocyclic Compounds (More Than One Hetero Atom)) Section cross-reference(s): 63, 78
- IT Musculoskeletal diseases
 - (tumor imaging; preparation of trihalobenzenetricarboxamide tristetraazacyclododecane metal complexes and related compds. as

```
contrast media)
    7429-91-6DP, Dysprosium, complexes 7439-89-6DP, Iron, complexes
     7439-96-5DP, Manganese, complexes 7440-53-1DP, Europium, complexes
     7440-54-2DP, Gadolinium, complexes 753020-30-3P 753020-31-4P
     753020-32-5P 753020-33-6P 753020-34-7P 753020-35-8P
     753020-36-9P 753020-37-0P 753020-39-2P 753020-40-5P 753020-42-7P 753020-43-8P 753020-44-9P 753020-45-0P 753020-46-1P
     753020-49-4P 753020-51-8P 753020-53-0P
     753020-56-3P 753020-59-6P 753020-61-0P
     753020-63-2P
                   753020-65-4P
     RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (preparation of trihalobenzenetricarboxamide tristetraazacyclododecane
metal
       complexes and related compds. as contrast media)
    14321-26-7P 88285-82-9P 138884-10-3P 138884-11-4P
ΙT
                                                            400894-66-8P
     425367-31-3P 425367-47-1P
                                  425367-60-8P
                                                752252-78-1P
                                                               752252-79-2P
                  752252-81-6P 752252-82-7P 752252-83-8P
     752252-80-5P
     752252-84-9P 752252-85-0P 752252-86-1P 752252-87-2P
     752252-88-3P 752252-89-4P
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                                                               752252-97-4P
     752252-98-5P 752252-99-6P 752253-00-2P 752253-01-3P 752253-02-4P
     752253-03-5P 752253-04-6P 752253-05-7P 752253-06-8P 752253-07-9P
     752253-08-0P 752253-09-1P 752253-10-4P 752253-11-5P
                                                               752253-12-6P
     752253-13-7P 752253-14-8P 752253-15-9P 752253-16-0P
                                                               752253-17-1P
     752253-18-2P 752253-19-3P 752253-20-6P 752253-21-7P
                                                               752253-22-8P
     752253-23-9P 752253-24-0P 752253-25-1P 752253-26-2P
     752253-27-3P 752253-28-4P 752253-29-5P 752253-30-8P
     752253-31-9P 752253-32-0P 752253-33-1P 752253-34-2P
     752253-35-3P 752253-36-4P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of trihalobenzenetricarboxamide tristetraazacyclododecane
metal
       complexes and related compds. as contrast media)
    753020-32-5P 753020-33-6P 753020-49-4P
     753020-51-8P 753020-53-0P 753020-56-3P
     753020-59-6P 753020-61-0P
     RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (preparation of trihalobenzenetricarboxamide tristetraazacyclododecane
metal
       complexes and related compds. as contrast media)
    753020-32-5 ZCAPLUS
RN
CN
    Yttrium, [u3-[[10,10',10''-[(2,4,6-triiodo-1,3,5-
     benzenetrivl)tris(carbonvlimino-2,1-ethanedivlimino(2-(oxo-κ0)-2,1-
     ethanediyl]]]tris[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-
     KN1, KN4, KN7, KN10, KO1, KO4, KO711(9-
     ) | | tri- (9CI) (CA INDEX NAME)
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10/573938

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PAGE 2-A

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PAGE 1-A

RN 753020-33-6 ZCAPLUS

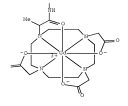
CN Gadolinium, [µ3-[[10,10',10''-[(2,4,5-triiodo-1,3,5-benzenetriyl)tris[carbonylimino-2,1-ethanediylimino[1-methyl-2-(oxo-KO)-2,1-ethanediyl]]]tris[[1,4,7,10-tetraazacyclododecane-1,4,7triacetato-kN1, kN4, kN7, kN10, kO1, kO4,.k appa.07][[9-]]tri-[(SCI) (CAINDEX NAME)

NH-CH2-CH2-NH-

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PAGE 2-B



RN 753020-49-4 ZCAPLUS

10/573938

- CN Gadolinium, [µ3-[[10,10'-[[2,4,6-triiodo-5-[[[[[4,7,10-tris[(carboxyk0)methyl]-1,4,7,10-tetraazacyclododec-1-ylkN1,kN4,kN7,kN10]acetyl
 - kO]amino]acety1]amino]-1,3-phenylene]bis[carbonylimino-2,1-ethanediyl]imino[2-(oxo-kO)-2,1-ethanediyl]]]bis[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-kN1,kN4,kN7,.kapp a.N10,kO1,kO4,kO7][9-)][tri-(9CI) (CA INDEX NAME)

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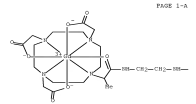
PAGE 2-B

RN 753020-51-8 ZCAPLUS

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phenylene|bis[carbonylimino-2,1-ethanediylimino[1-methyl-2-(oxo-KO)-2,1-ethanediyl]]]bis[1,4,7,10-tetraazacyclododecane-1,4,7-triacetatokN1,kN4,kN7,kN10,kO1,KO4,KO7][9-

)]]tri- (9CI) (CA INDEX NAME)



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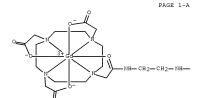
10/573938

PAGE 2-B

RN 753020-53-0 ZCAPLUS

CN Gadolinium, [µ3-[[10,10'-[[2,4,6-triiodo-5-[methyl[[[[4,7,10-tris([carboxy-KO]methyl]-1,4,7,10-tetraazacyclododec-1-yl-KNI, xN4, kN7, kN10] acetvl-

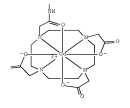
k0]amino]acety1]amino]-1,3-phenylene]bis[carbonylimino-2,1-ethanediyllimino[2-(oxo-k0)-2,1-ethanediyl]]]bis[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-kN1,kN4,kN7,.kapp a.N10,k01,k04,k07][9-)][tri-(9CI) (CA INDEX NAME)



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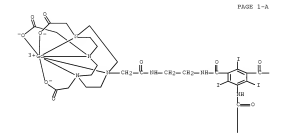


RN 753020-56-3 ZCAPLUS

10/573938

CN Gadolinium, [µ3-[[10,10'-[[2,4,6-triiodo-5-[[[[2-[[4,7,10-tris[(carboxy-K0)methyl]-1,4,7,10-tetraazacyclododec-1-yl-KN1,KN4,KN7,KN10]acetyl]amino]ethyl]amino]carbonyl

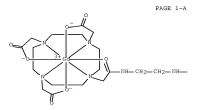
[amino]-1,3-phenylene]bis[carbonylimino-2,1-ethanediylimino[2-(oxo-K0)-2,1-ethanediyl]]]bis[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-KN1,KN4,KN7,KN10,K01,K04,LK
appa.07]](9-)]tri-(9C1) (CA INDEX NAME)



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RN 753020-59-6 ZCAPLUS

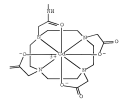
CN Gadolinium, [µ3-[10,10',10''-[(2,4,6-triiodo-1,3,5-benzenetriyl)tris[carbonylimino-2,1-ethanediylimino[2-(oxo-KO)-2,1-ethanediyl]]]tris[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-KNl,KN4,KN7,KN10,KOl,KO4,KO7]][9)]]tri-[9CI) (CA INDEX NAME)



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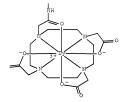


- RN 753020-61-0 ZCAPLUS
- CN Dysprosium, [μ3-[[10,10',10''-[(2,4,6-triiodo-1,3,5-benzenetriy])tris[carbonylimino-2,1-ethanediylimino[2-(oxo-κ0)-2,1-ethanediyl]]]tris[[1,4,7,10-tetraazacyclododecane-1,4,7-triacetato-κΝ],κΝθ,κΝ1,κΝ10,κΝ10,κΝ10,κΝ1,κΝή,κΝ9]
 -)]]tri- (9CI) (CA INDEX NAME)

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PAGE 2-B



752252-82-7P 752252-85-0P 752253-24-0P ΙT 752253-27-3P 752253-32-0P 752253-36-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of trihalobenzenetricarboxamide tristetraazacyclododecane metal

complexes and related compds. as contrast media)

- RN 752252-82-7 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10',10''-[(2,4,6triiodo-1,3,5-benzenetriyl)tris[carbonylimino-2,1-ethanediylimino(2-oxo-2,1-ethanedivl)]|tris- (9CI) (CA INDEX NAME)

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-- CH2-CO2H

103

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- RN 752252-85-0 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10',10''-[(2,4,6-triodo-1,3,5-benzenetriyl)|tris|carbonylimino-2,1-ethanediyl|mino(1-methyl-2-oxo-2,1-ethanediyl)||tris-(901) (CA INDEX NAME)

-- СН2-СО2Н

RN 752253-24-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10'-[(2,4,6-triiodo-5-[[[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]amino]acetyl]amino]-1,3-phenylene)bis[carbonylimino-2,1-ethanediylimino(2-oxo-2,1-ethanediyl)]bis-(9CI) (CA INDEX NAME)

PAGE 1-B

-CH2-CO2H

RN 752253-27-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10'-[(2,4,6-triodo-5-[[[[1-cxo-2-[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-y1]propyl]amino]acetyl]amino]-1,3-phenylene)bis[carbonylimino-2,1-ethanediylimino(1-methyl-2-oxo-2,1-ethanediyl)]bis-(9CI) (CA INDEX NAME)

$$- \text{CH}_2 - \text{CH}_2 - \text{NH} - \text{CH}_2 - \text{CO}_2 H$$

$$- \text{CH}_2 - \text{CH}_2 - \text{NH} - \text{CH}_2 - \text{CO}_2 H$$

$$- \text{CH}_2 - \text{CO}_2 H$$

PAGE 2-B

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RN 752253-32-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10'-[(2,4,6-triiodo-5-[methyl[[(4,7,10-tria(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]amino]acetyl]amino]-1,3-phenylene)bis[carbonylimino-2,1-ethanediylimino(2-oxo-2,1-ethanediyl)]bis-(9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{PAGE 1-A} \\ \text{HO}_2\text{C}-\text{CH}_2 \\ \text{HO}_2\text{C}-\text{CH}_2 \\ \end{array} \\ \text{N} \\ \text{HO}_2\text{C}-\text{CH}_2 \\ \text{N} \\ \text{HO}_2\text{C}-\text{CH}_2 \\ \end{array}$$

$$\begin{array}{c} \text{HO}_2\text{C}-\text{CH}_2\\ \text{HO}_2\text{C}-\text{CH}_2\\ \text{HO}_2\text{C}-\text{CH}_2 \end{array} \\ \text{N} \qquad \text{CH}_2-\text{C}-\text{NH}-\text{CH}_2-\text{CH}_2-\text{NH}-\text{CH}_2\\ \text{HO}_2\text{C}-\text{CH}_2\\ \end{array}$$

RN 752253-36-4 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10'-[(2,4,6-triodo-5-[[[2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]amino]ethyl]amino]carbonyl]amino]-1,3-phenylene)bis[carbonylimino-2,1-ethanediylimino(2-oxo-2,1-ethanediyl)]]bis-(9CI) (CA INDEX NAME)

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PAGE 2-B

-CH2-CO2H

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 10 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2004:432097 ZCAPLUS Full-text

DOCUMENT NUMBER: 141:153123

TITLE: In Vitro and in Vivo Comparison of Human Escherichia coli Heat-Stable Peptide Analogues Incorporating the

111In-DOTA Group and Distinct Linker Moieties
AUTHOR(S): Giblin, Michael F.; Gali, Hariprasad; Sieckman, Gary

L.; Owen, Nellie K.; Hoffman, Timothy J.; Forte,

Leonard R.; Volkert, Wynn A.

CORPORATE SOURCE: Research Service, Harry S. Truman Memorial Veterans'
Administration Hospital, Columbia, MO, 65201, USA

SOURCE: Bioconjugate Chemistry (2004), 15(4), 872-880 CODEN: BCCHES; ISSN: 1043-1802

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Three human Escherichia coli heat-stable peptide (STh) analogs, each containing a DOTA chelating group, were synthesized by SPPS and oxidative refolding and compared in in vitro and in vivo systems. One analog, DOTA-F19-STh(1-19), contains an N-terminal DOTA group attached via an amide bond linkage to an STh moiety which is essentially wild-type except for a Tyr to Phe alteration at position 19 of the mol. A second analog, DOTA-R1, 4, F19-STh(1-19), differs from the first in that asparagine residues in positions 1 and 4 have been altered to arginine residues in order to examine the effect of pos. charged groups in the linker domain. A third analog, DOTA-11AUN-F19-STh(1-19), differs from the first in that it incorporates an 11aminoundecanoic acid spacer group between the DOTA group and the first asparagine residue. In vitro competitive binding assays utilizing T-84 human colon cancer cells demonstrated that significant alterations to the N-terminal region of the STh mol. were well tolerated and did not significantly affect binding affinity of STh for the guanylyl cyclase C (GC-C) receptor.

Internalization and efflux studies of the indium-labeled species demonstrated that inclusion of pos. charge in the linker moiety inhibits internalization of the compound within tumor cells. The characteristics of the three analogs were compared in an in vivo model utilizing T-84 human colon cancer cell xenografts in SCID mice. Clearance of all analogs was rapid, primarily via renal excretion into the urine, with >89% ID excreted into the urine at 1 h pi for all analogs. The 111In-DOTA-R1, 4, F19-STh(1-19) and 111In-DOTA-11AUN-F19-STh(1-19) analogs both had longer residence times in the blood than did the 111In-DOTA-F19-STh(1-19) analog, probably accounting for increased %ID/g values for tumors and nontarget tissues at 1 h pi. At 4 h pi, significant differences between analogs were only seen with respect to metabolic routes of excretion, indicating that increased blood residence time did not result in increased tumor residualization. Reduction of hepatic uptake of these compds., however, could have significance in the development of agents for the imaging of hepatic metastases. The ability to manipulate in vivo pharmacodynamics and tumor uptake of radiolabeled STh peptides through modification of linker moieties is under continuing investigation in order to produce optimal imaging and therapeutic radiopharmaceuticals.

CC 8-9 (Radiation Biochemistry) IT 415706-07-9P 723914-72-5P 728914-74-7P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (in vitro and in vivo comparison of human E. coli heat-stable peptide

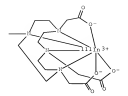
analogs incorporating 111In-DOTA group and distinct linker moieties)

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (in vitro and in vivo comparison of human E. coli heat-stable peptide analogs incorporating 1111n-DOTA group and distinct linker moieties)

415706-07-9 ZCAPLUS

RN

CN Indate(2-)-111In, [N2-[2-[4,7,10-tris](carboxy-kO)methyl]-1,4,7,10-tttaazacyclododec-1-yl-kNl,kNl,kNl,kNl)acetyl]-L-asparaginyl-L-seryl-L-seryl-L-asparaginyl-L-tyrosyl-L-cysteinyl-L-cysteinyl-L-cysteinyl-L-aphanyl-L-l-cysteinyl-L-tyrosyl-L-cysteinyl-L-aphanyl-L-cysteinyl-L-threonylglycyl-L-cysteinyl-L-phenylalanine cyclic (6-tl),(7-tl),(10-tl)-tris(disulfidato)(5-)]-, hydrogen (1:2) (CA INDEX NAME)



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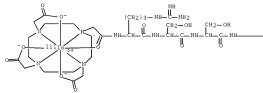
Ph-CH2-

●2 H+

- RN 728914-72-5 ZCAPLUS
- CN Indate(2-)-1111n, [N2-[[4,7,10-tris[(carboxy- κ 0)methy1]-1,4,7,10-tetraazacyclododec-1-yl- κ N1, κ N4, κ N7, κ N10]acetyl-

$$\begin{split} \kappa O] = & L - arginyl - L - seryl - L - aerylnyl - L - tyrosyl - L - cysteinyl - C - cysteinyl - L - cysteinyl - C - cystei$$





PAGE 1-B

PAGE 2-A

PAGE 3-B

— Ph

RN 728914-74-7 ZCAPLUS

CN Indate(2-)-1111n, [N2-[1-oxo-11-[[[4,7,10-tris[(carboxy- κ 0)methy1]-1,4,7,10-tetraazacyclododec-1-y1- κ N1, κ N4, κ N7, κ N10]

PAGE 1-B

PAGE 4-B

— Ph

PAGE 5-A

REFERENCE COUNT:

THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 11 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2004:261461 ZCAPLUS Full-text

49

DOCUMENT NUMBER: 142:70820

TITLE: AUTHOR(S):

Cellular Delivery of MRI Contrast Agents Allen, Matthew J.; MacRenaris, Keith W.; Venkatasubramanian, P. N.; Meade, Thomas J.

Dep. Chem., Biochem. and Mol. and Cell Biol., CORPORATE SOURCE: Neurobiol. and Physiol., and Radiol., Northwestern

Univ., Evanston, IL, 60208, USA SOURCE: Chemistry & Biology (2004), 11(3), 301-307

CODEN: CBOLE2: ISSN: 1074-5521

PUBLISHER: Cell Press DOCUMENT TYPE: Journal LANGUAGE: English

Magnetic resonance imaging (MRI) is a powerful tool for acquiring images of opaque living animals with the benefit of tracking events over extended periods of time on the same specimen. Contrast agents are used to enhance regions, tissues, and cells that are magnetically similar but histol. distinct. A principal barrier to the development of MRI contrast agents for investigating biol. questions is the delivery of agents across cellular membranes. Here, we describe the synthesis and in vitro testing of Gd(III)based MRI contrast agents containing varying length polyarginine oligomers capable of permeating cell membranes. We examine the effect of the length of oligomer on T1 enhancement and cellular uptake. Furthermore, the effect of incubation time, concentration, and cell type on uptake is explored. Toxicity and washout studies are performed in addition to MRI phantom studies.

8-9 (Radiation Biochemistry)

22541-18-0DP, Europium III, complexes with DOTA-polyarginine, biological studies 22541-19-1DF, Gadolinium(III), complexes with DOTA-polyarginine, biological studies 812644-18-1P 812644-19-2P 812644-20-5P 812644-21-6P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Gd(III)-based MRI contrast agents preparation and cellular uptake) ΤТ 811804-40-7P 811804-47-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

- (Gd(III)-based MRI contrast agents preparation and cellular uptake)
 IT 22541-138-0PP. Ruropium III, complexes with DOTA-polyarginine,
 biological studies 22541-19-1DP, Gadolinium(III), complexes with
 DOTA-polyarginine, biological studies 812644-18-1P
 81264-19-2P 812644-20-5P 812644-21-6P
 RE: DGM (Diagnostic use); PRT (Pharmacokinetics); PRP (Properties); SPN
 (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES
 (Uses)
 (Gd(III)-based MRI contrast agents preparation and cellular uptake)
- (GG(III)-based MRI contrast agents preparation and cellular uptake RN 22541-18-0 ZCAPLUS
- CN Europium, ion (Eu3+) (CA INDEX NAME)

Eu3+

- RN 22541-19-1 ZCAPLUS
- CN Gadolinium, ion (Gd3+) (CA INDEX NAME)

Gd 3+

- RN 812644-18-1 ZCAPLUS
- CN Gadolinate(1-), [N2-[[4,7,10-tris](carboxy-KO)methyl]-1,4,7,10-tetraazacyclododec-l-yl-kNl, kN4,kN7,kNl0]acetylKO]-L-arginyl-L-argi

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PAGE 1-B

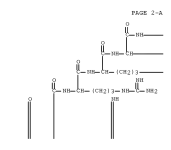
PAGE 2-A

H+

- RN 812644-19-2 ZCAPLUS
- CN Gadolinate(1-), [N2-[[4,7,10-tris](carboxy-k0)methyl]-1,4,7,10tetrazacyclododec-l-yl-kNl,kN4,kN7,kNl0]acetylk0]-L-arginyl-L-

PAGE 1-B

PAGE 2-B



CH_ (CH2)3_NH_C_NH2 __(CH₂)₃_NH_C_NH₂

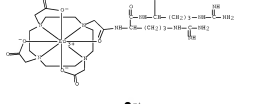
- RN 812644-20-5 ZCAPLUS
- CN Europate(1-), [N2-[[4,7,10-tris[(carboxy-KO)methyl]-1,4,7,10-tetrazacyclododec-1-yl-kNl, kN4,kN7,kNl0]acetyl-KO]-L-arginyl-L-arginy

PAGE 1-A

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PAGE 1-B

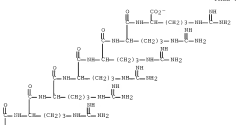
PAGE 2-A

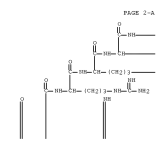


RN 812644-21-6 ZCAPLUS

PAGE 1-B

PAGE 2-B





- IT 811804-40-7P 811804-47-4P
- RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
- (Gd(III)-based MRI contrast agents preparation and cellular uptake) RN 811804-40-7 ZCAPLUS
- CN L-Arginine, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-arg

Absolute stereochemistry.

RN 811804-47-4 ZCAPLUS

CN L-Arginine, N2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-arg

Absolute stereochemistry.

PAGE 2-A

PAGE 3-A

REFERENCE COUNT: 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 12 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2003:750102 ZCAPLUS Full-text

DOCUMENT NUMBER: 139:214227

TITLE: A New Biotin Derivative-DOTA Conjugate as a Candidate for Pretarqued Diagnosis and Therapy of Tumors AUTHOR(S): Sabatino, Giuseppina; Chinol, Marco; Paqanelli,

Giovanni; Papi, Stefano; Chelli, Mario; Leone, Giuseppe; Papini, Anna Maria; De Luca, Angelo;

Ginanneschi, Mauro

CORPORATE SOURCE: Dep. of Org. Chem. "Ugo Schiff", CNR-ICCOM, Polo Scientifico, Univ. of Florence, Sesto Fiorentino, I-50019, Italy

SOURCE: Journal of Medicinal Chemistry (2003), 46(14), 3170-3173

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society
DOCUMENT TYPE: Journal

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:214227

- AB The synthesis of a new biotin derivative, the (CO) reduced N-aminohexyl biotinamido derivative, designed to be serum biotinidase resistant, and its conjugation to the chelator DOTA through an amide bond at one of the four carboxymethyl chains are described. The 90Y-labeled conjugate was able to bind avidin at different Av/conjugate molar ratios with good results. The preclin. The preclin. results indicate that this new biotin-DOTA conjugate is a good candidate for pretargeted diagnosis and therapy of tumors.
- CC 26-8 (Biomolecules and Their Synthetic Analogs) Section cross-reference(s): 1
- ST aminohexyl biotinamido biotin deriv prepn; biotin DOTA conjugate prepn; stability avidin binding biotin DOTA conjugate; pretargeted diagnosis tumor therapy biotin DOTA conjugate prepn
- IT Antitumor agents

Diagnostic agents

(preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)

- IT Avidins
 - RL: BSU (Biological study, unclassified); BIOL (Biological study) (preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)
- IT 451478-45-8P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)

- IT 586962-90-5P
 - RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
 - (preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)
- IT 58-85-5 51857-17-1 60239-18-1
 - RL: RCT (Reactant); RACT (Reactant or reagent)
 - (preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)
- IT 65953-56-2P 153162-70-0P 451478-44-7P
 - RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 - (preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)
- IT 451478-45-8P
 - RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)
 - (preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)
- RN 451478-45-8 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4
 - vl|pentvl|amino|hexvl|amino|-2-oxoethvl|- (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

__CO2H

IT 586962-90-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of a biotin-DOTA conjugate and evaluation of its serum stability and avidin binding properties as a candidate for pretargeted diagnosis and potential tumor therapy)

RN 586962-90-5 ZCAPLUS

CN Yttrium-90Y, [10-[2-[[6-[[5-(hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl)pentyl]amino]hexyl]amino]-2-(oxo-ko)ethyl]-1,4,7,10-terrazacyclododecane-1,4,7-triacetato(3-)-kNl,kNl,kNl7,.kappa.Nl0,kOl,ko4,kO7]-(9CI) (CA INDEX NAME)

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 13 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2003:726127 ZCAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 140:299530

TITLE: Synthesis and visualization of a membrane-permeable

MRI contrast agent
AUTHOR(S): Allen, Matthew J.;

AUTHOR(S): Allen, Matthew J.; Meade, Thomas J.

CORPORATE SOURCE: Division of Biology and the Beckman

URCE: Division of Biology and the Beckman Institute, California Institute of Technology, Pasadena, CA, 91125, USA

SOURCE: JBIC, Journal of Biological Inorganic Chemistry

(2003), 8(7), 746-750

CODEN: JJBCFA; ISSN: 0949-8257

PUBLISHER: Springer-Verlag

DOCUMENT TYPE: Journal LANGUAGE: English

AB The study of in vivo developmental events has undergone significant advances with the advent of biol, mol, imaging techniques such as computer enhanced light microscopy imaging, positron emission tomog. (PET), micro-CT, and magnetic resonance imaging (MRI). MRI has proven to be a particularly powerful tool in clin. and biol. settings. Images can be acquired of opaque living animals, with the benefit of tracking events of extended periods of time on the same specimen. Contrast agents are routinely used to enhance regions, tissues, and cells that are magnetically similar but histol. distinct. A principal barrier to the development of MR contrast agents for investigating developmental biol. questions is the ability to deliver the agent across cellular membranes. As part of our research, we are investigating a number of small mols. that facilitate transport of charged and uncharged species across cell membranes. Here we describe the synthesis and testing of a Gd(III)-based MR contrast agent conjugated to polyarginine that is able to permeate cell membranes. We confirmed cellular uptake of the agent using twophoton laser microscopy to visualize a Eu(III) derivative of the contrast agent in cell culture, and verified this uptake by Tl anal. of the Gd(III) agent in cells.

8-9 (Radiation Biochemistry)

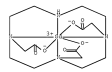
IT 112188-16-6P 137184-55-5P

RL: PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

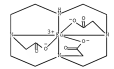
(preparation of Gd(III)-based membrane-permeable MRI contrast agent)
T 576553-18-78 676553-19-88

RL: PKT (Pharmacokinetics); SPN (Synthetic preparation); THU (Therapeutic

- use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of Gd(III)-based membrane-permeable MRI contrast agent)
- RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP
 (Preparation); RACT (Reactant or reagent)
- (preparation of Gd(III)-based membrane-permeable MRI contrast agent)
 IT 7087-68-5, Diisopropylethylamine 91000-69-0D, L-Arginine,
- N2-[(9H-fluoren-9-ylmethoxy)carbonyl]-, resin-bound 137076-54-1, DOTA tri(tert-butyl) ester 148893-10-1, HATU 6/8544-85-7 RL: RCT (Reactant); RACT (Reactant or reagent)
- (preparation of Gd(III)-based membrane-permeable MRI contrast agent)
 II 112188-16-6P 137184-55-5P
 - RL: PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
- (preparation of Gd(III)-based membrane-permeable MRI contrast agent)
 RN 112188-16-6 ZCAPLUS
- CN Gadolinium, [1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)KN1,KN4,KN7,KN10,KO1,KO4,KO7](9C1) (CA INDEX NAME)



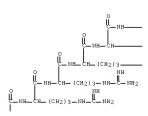
- RN 137184-55-5 ZCAPLUS
- CN Europium, [1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)kNl,kN4,kN7,kNl0,kOl,kO4,kO7]-(9C1) (CA INBEX NAME)



- IT 676553-18-7P 676553-19-8P
 - RL: PKT (Pharmacokinetics); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation) of Gd(III)-based membrane-permeable MRI contrast agent)
- RN 676553-18-7 ZCAPLUS

CN Gadolinate(1-), [N2-[[4,7,10-tris](carboxy-k0)methyl]-1,4,7,10-tetrazacyclododec-1-y1-kN1, kN4, kN7, kN10]acetyl]-L-arginyl

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PAGE 2-A

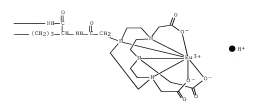
PAGE 3-B

- RN 676553-19-8 ZCAPLUS
- CN Europate(1-), [N2-[[4,7,10-tris[(carboxy-KO)methyl]-1,4,7,10tetraazacyclododec-1-y1-KN1, KN4, KN7, KN10]acetyl]-Larginyl-L-a

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- IT 676544-84-6P
- RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of Gd(III)-based membrane-permeable MRI contrast agent)
- RN 676544-84-6 ZCAPLUS
- CN L-Arginine, N2-[[4,7,10-tris(carboxymethy1)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-L-arginyl-L-arg

Absolute stereochemistry.

IT 676544-85-7

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of Gd(III)-based membrane-permeable MRI contrast agent)

RN 676544-85-7 ZCAPLUS

CN Europium hydroxide (Eu(OH)3), pentahydrate (9CI) (CA INDEX NAME)

HO_Eu_OH

●5 H2O

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 14 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2003:71732 ZCAPLUS Full-text

ACCESSION NUMBER: 2003:71732 DOCUMENT NUMBER: 138:122864

TITLE: Preparation of vitronectin receptor antagonist

pharmaceuticals for use in the diagnosis and treatment

INVENTOR(S): Harris, Thomas D.; Barrett, John A.; Carpenter, Alan

P., Jr.; Rajopadhye, Milind

P., Jr.; Kajopadnye, Milind PATENT ASSIGNEE(S): USA

SOURCE: U.S., 146 pp., Cont.-in-part of U.S. Ser. No. 465,300.

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE.

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 8

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6511649	B1	20030128	US 2000-599364	20000621

US	6322770	B1	20011127	US	1999-281207		19990330
US	2002015680	A1	20020207	US	1999-281209		19990330
US	6524553	B2	20030225				
US	6548663	B1	20030415	US	1999-281050		19990330
US	2002182147	A1	20021205	US	1999-465300		19991217
US	6511648	B2	20030128				
US	2002041878	A1	20020411	US	2001-948807		20010907
US	6683163	B2	20040127				
US	2002061909	A1	20020523	US	2001-948390		20010907
US	6689337	B2	20040210				
US	2003232053	A1	20031218	US	2001-947783		20010907
US	6743412	B2	20040601				
US	2003124120	A1	20030703	US	2002-269252		20021011
US	2003113336	A1	20030619	US	2002-281015		20021026
US	7018611	B2	20060328				
US	2003149262	A1	20030807	US	2002-306054		20021126
PRIORITY	APPLN. INFO.:			US	1998-112732P	P	19981218
				US	1999-465300	A2	19991217
				US	1998-80150P	P	19980331
				US	1998-112715P	P	19981218
				US	1998-112829P	P	19981218
				US	1998-112831P	P	19981218
					1999-281050		19990330
				US	1999-281209	A3	19990330
				US	2000-599364	A3	20000621

OTHER SOURCE(S): MARPAT 138:122864

AB Compds. (Q)d-Ln-Ch and (Q)d-Ln-(Ch)d' [Q is a residue having a quinolone-type modety; In is a linking group; Ch is a metal-bonding unit; d = 1-10; d' = 1-100] and pharmaceutical compns. containing them were prepared for the treatment of cancer in combination therapy. The pharmaceuticals are comprised of a targeting modety that binds to a receptor that is upregulated during angiogenesis, an optional linking group, and a therapeutically effective radioisotope or diagnostically effective imageable modety. The imageable modety is a gamma ray or positron emitting radioisotope, a magnetic resonance imaging contrast agent, an X-ray contrast agent, or an ultrasound contrast agent. Thus, 2-[[[4-[4-[(]3-[2-[2-[3-[[6-[[1-aza-2-(2-sufoobenvil)vivil]aminol-3-

pyridyl]carbonylamino]propoxy]ethoxy]ethoxy]propyl]amino]sulfonyl]phenyl]phenyl]sulfonyl]amino]-3-[[7-[(imidazo1-2-ylamino)methyl]-1-methyl-4-oxo-3-hydroquinolyl]carbonylamino]propanoic acid (claimed compound) was prepared ICM A61K051-00

ICS A61M036-14

INCL 424001690; 424001110; 424001650; 424009100; 424009400; 424009500; 530331000

CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 8, 27, 28, 63, 78

IT Angiogenesis

Antirheumatic agents

Antitumor agents

Human

numan

Imaging agents

Radiopharmaceuticals

(preparation of peptide- and tetraazadodecane-containing quinolones and their

radioactive metal complexes for diagnosis and treatment of cancer) 17 5704-04-1DP, Tricine, technetium-99 complexes 10098-91-6DP, complexes with vitronectin receptor binding conjugates, preparation 14133-76-7DP, complexes with vitronectin receptor binding conjugates, preparation 14265-75-9DP, complexes with vitronectin receptor binding conjugates, preparation 15750-15-9DP, complexes with vitronectin receptor binding

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conjugates, preparation 63995-70-0DP, TPPTS, technetium-99 complexes
277315-51-2P 277315-52-3P 277315-53-4P 277315-55-6P 277315-56-7P
277315-57-8P 277315-58-9P
                           277315-59-0P 277315-60-3P
                                                      277315-61-4P
277315-62-5P 277315-63-6P 277315-64-7P 277315-65-8P
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                                        277315-72-7P
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277315-78-3P
            277315-79-4P
                          277315-80-7P 277315-81-8DP, technetium-99
complexes 277316-60-6P 277316-61-7P 277316-62-8P 277316-63-9P
277316-64-0P 277316-65-1P 277316-66-2P 277316-67-3P
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278173-00-5P
             278173-01-6P 278173-02-7P
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278173-04-9P
            278173-05-0P
                           278173-06-1P
                                         278173-07-2P
278173-08-3P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
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(preparation of peptide- and tetraazadodecane-containing quinolones and

their

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radioactive metal complexes for diagnosis and treatment of cancer)
тт
    40324-66-1P 57932-18-0P
                             137076-54-1P
                                          192635-89-5P 220156-99-0P
    250612-31-8P 277315-82-9P
                               277315-83-0P
                                            277315-84-1P 277315-85-2P
    277315-86-3P 277315-88-5P 277315-89-6P 277315-90-9P
                                                          277315-91-0P
    277315-92-1P 277315-93-2P 277315-94-3P 277315-95-4P 277315-96-5P
    277315-97-6P 277315-98-7P 277315-99-8P 277316-00-4P 277316-01-5P
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                              277316-06-0P
                                            277316-08-2P
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    277316-39-9P 277316-40-2P
                              277316-41-3P
                                            277316-42-4P 277316-43-5P
    277316-44-6P
                 277316-45-7P 277316-47-9P 277316-48-0P
    277316-50-4P
                 277316-52-6P 277316-53-7P 277316-54-8P 277316-56-0P
    277316-58-2P
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

 $\hbox{ (preparation of peptide- and tetraazadodecane-containing quinolones and their}\\$

radioactive metal complexes for diagnosis and treatment of cancer) IT -277315-74-9P 277315-75-0P 278173-04-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of peptide- and tetraazadodecane-containing quinolones and

their

radioactive metal complexes for diagnosis and treatment of cancer)

RN 277315-74-9 ZCAPLUS
1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[[(2S)-2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

RN 277315-75-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[[(2S)-2-carboxy-2-[([2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2-oxoethyl]-, tris(trifluoroacetate) (9CI) (CA INDEX NAME)

CM :

CRN 277315-74-9

CMF C45 H61 N11 O13 S

Absolute stereochemistry.

PAGE 1-A

CM 2 CRN 76-05-1 CMF C2 H F3 O2

RN 278173-04-9 ZCAPLUS

N Yttrate(1)-90%, [10-[2-[[3-[3-[[[(28)-2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino]methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2-(oxo-x0)ethyl]-1, 4, 7, 10-tetraazacyclododecane-1, 4, 7-triacetato(4-)- kN1, kN4, kN7, kN10, kO1, kO4, kO7)-, hydrosen (9CI) (CA INDEX NAME)

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IT 277316-47-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

 $\label{preparation} \mbox{ (preparation of peptide- and tetraazadodecane-containing quinolones and their}$

radioactive metal complexes for diagnosis and treatment of cancer)

- RN 277316-47-9 ZCAPLUS CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, $10-[2-[[3-[3-[[(2S)-2-carboxy-2-[(2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2-oxoethyl]-, <math>\alpha,\alpha',\alpha''$ -tris(1,1-dimethylethyl) ester, tris(trifluoroacetate) (SCI) (Δ A INDEX NAME)
 - CM 1
 - CRN 277316-46-8
 - CMF C57 H85 N11 O13 S

Absolute stereochemistry.

PAGE 1-A

CM 2

CRN 76-05-1 CMF C2 H F3 O2

F

REFERENCE COUNT:

143 THERE ARE 143 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 15 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2001:935440 ZCAPLUS <u>Full-text</u> DOCCUMENT NUMBER: 136:70082

TITLE:

Vitronectin receptor antagonist pharmaceuticals for

10/573938 INVENTOR(S):

use in combination therapy

Harris, Thomas D.; Barrett, John A.; Carpenter, Alan

P., Jr.; Rajopadhye, Milind

PATENT ASSIGNEE(S): Dupont Pharmaceuticals Company, USA; Bristol-Myers Squibb Pharma, Company

PCT Int. Appl., 542 pp.

SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE:

English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					DATE		APPLICATION NO.											
WO				A2 20011227		WO 2001-US19793													
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		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP	, 1	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	
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		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML	, 1	MR,	NE,	SN,	TD,	TG			
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BR	2001	0118	80		A		2006	0425		BR	20	01-	1188	0		2	0010	621	
NZ	5229	25			A		2006	0831		NZ	20	01-5	5229	25		2	0010	621	
MX	MX 2002PA12750 A 20			20040730 MX 2002-PA1275				750	20021218										
IN	2007	DN01	157		A		2007	0427		IN	20	07-1	ON11.	57		2	0070	213	
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OTHER SOURCE(S): MARPAT 136:70082

Anticancer agents of the formulas (Q)d-Ln-Ch or (Q)d-Ln-(Ch)d (I) [Q is a residue having a quinolone-type moiety; Ln is a linking group; Ch is a metalbonding unit; d = 1-10; d' = 1-100] and kits containing I are prepared for the treatment of cancer in combination therapy in a patient. I are comprised of a targeting moiety that binds to a receptor that is upregulated during angiogenesis, an optional linking group, and a therapeutically effective radioisotope or diagnostically effective imageable moiety. I may be used with radioisotopes; in addition, I may be used in conjunction with radio- and photosensitizers, ligands such as TPPTS or tricine, and reducing agents such as tin(II). The present invention provides novel compds. useful for the treatment of rheumatoid arthritis (no data).

- IC. ICM A61K041-00 TCS A61K051-04
- 34-3 (Amino Acids, Peptides, and Proteins)
- Section cross-reference(s): 8, 27, 28, 63, 78
- Antitomor agents

(preparation of peptide- and tetraazadodecane-containing quinolones and

their

- radioactive metal complexes as anticancer agents)
- 5704-04-1DP, Tricine, technetium-99 complexes 10098-91-6DP, complexes with vitronectin receptor binding conjugates, preparation 14133-76-7DP,

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complexes with vitronectin receptor binding conjugates, preparation
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preparation 15750-15-9DP, complexes with vitronectin receptor binding
conjugates, preparation 63995-70-0DP, TPPTS, technetium-99 complexes
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             277315-58-9P 277315-59-0P
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278173-04-9P 278173-05-0P 278173-06-1P 278173-07-2P
278173-08-3P
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(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (preparation of peptide- and tetraazadodecane-containing quinolones and
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their

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radioactive metal complexes as anticancer agents)
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    277316-39-9P 277316-40-2P
                              277316-41-3P
                                            277316-42-4P
                                                          277316-43-5P
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of peptide- and tetraazadodecane-containing quinolones and

their

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radioactive metal complexes as anticancer agents)
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277315-74-9P 277315-75-0P 278173-04-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of peptide- and tetraazadodecane-containing quinolones and

their

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radioactive metal complexes as anticancer agents)
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RN 277315-74-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[[(2S)-2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2oxoethvll- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

RN 277315-75-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[[(28)-2-carboxy-2-[([2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2-oxoethyl]-, tris(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 277315-74-9

CMF C45 H61 N11 O13 S

PAGE 1-A

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 278173-04-9 ZCAPLUS

CN Yttrate(1-)-90%, [10-[2-[[3-[3-[1](2S)-2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]mino]ethyl]mino]carbonyl]-7-[(1H-imidazol-2-ylmino)methyl]-4-oxo-1(4H)-quinollinyl]propyl]mino]-2-[oxo-k0]ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(4-)-kN1,kN4,kN7,kN10,kO1,kO4,kO7]-,hydrogen (9CI) (CA INDEX NAME)

PAGE 2-A

IT 277316-47-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

 $\label{preparation} \mbox{ (preparation of peptide- and tetraazadodecane-containing quinolones and their}$

radioactive metal complexes as anticancer agents)

tris(trifluoroacetate) (9CI) (CA INDEX NAME)

CM :

CRN 277316-46-8

CMF C57 H85 N11 O13 S

PAGE 1-A

PAGE 2-A со₂н о

CM 2

CRN 76-05-1 CMF C2 H F3 O2

L79 ANSWER 16 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:661180 ZCAPLUS Full-text DOCUMENT NUMBER: 133:249059

TITLE:

Radionuclide conjugates with DOTA-biotin derivatives for diagnosis and therapy INVENTOR(S): Griffiths, Gary L.; Hansen, Hans; Govindan, Serengulam

148

DOCUMENT TYPE:

PATENT ASSIGNEE(S): Immunomedics, Inc., USA

SOURCE: U.S., 10 pp., Cont.-in-part of U.S. Ser. No. 486,166, abandoned.

CODEN: USXXAM Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 19

PATENT INFORMATION:

PATENT NO.				KIN	D	DATE			APPLICATION NO.					DATE					
US 6120768			A 20000919				US 1997-990843												
US 5736119			A	A 19980407				US 1	995-	4099	19950323								
US	US 5922302			A		1999	0713	US 1995-440652						19950515					
WO	9930	745			A2	A2 19990624				WO 1998-US26579					19981215				
WO	WO 9930745				A3 20000113														
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		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	UA,	UG,		
		US,	UZ,	VN,	YU,	ZW													
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		CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG								
AU	AU 9918258				A	A 19990705				AU 1999-18258					19981215				
PRIORIT	Y APP	LN.	INFO	. :						US 1993-62662				B1 19930517					
							US 1995-409960												
							US 1995-486166												
							US 1996-688781												
										US 1						9971			
										WO 1						9981			
										MO T	フフロー	0526	219		M T	フフロエ	Z13		

AB A radionuclide-chelator conjugate composition for detecting and/or treating lesions in a patient comprises pre-targeting the cell, tissue, or pathogen with a substrate, using a targeting protein that specifically binds a marker substance on the target cell, tissue, or pathogen and to which the substrate is directly or indirectly bound. Parenteral injection comprises a chelate conjugate of biotin, a chelator, and a chelatable detection or therapeutic agent, and allows the composition to accrete at the targeted cell, tissue, or pathogen. The chelate conjugate is purified by liquid chromatog. after chelate formation, or further comprises a blood transit-modifying linker or addend that is covalently bound within the chelate conjugate, or both. The detection or therapeutic agent of the invention are used to detect or treat cancer, infectious diseases, or cardiovascular diseases. Preparation of biotin-D-Ph-D-Jys-DOTA is presented.

IC ICM A61K039-395

INCL 424178100

CC 8-9 (Radiation Biochemistry)

Section cross-reference(s): 1, 15, 28, 34

ST DOTA biotin deriv chelator radionuclide conjugate; diagnosis therapy DOTA biotin deriv radionuclide; antitumor antiinfective cardiovascular agent radionuclide conjugate

IT Antitumor agents

(carcinoma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

IT Antitomor agents

(glioma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

IT Antitumor agents

(leukemia; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy) $\,$

IT Antitumor agents

(lymphoma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

IT Antitumor agents

(melanoma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

IT Antitumor agents

(myeloma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

IT Antitumor agents

(neuroblastoma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

IT Anti-infective agents

Antimicrobial agents

Antitumor agents

Cardiovascular agents

Parasiticides

 $\mbox{(radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)}$

Antitumor agents

(sarcoma; radionuclide conjugates with DOTA-biotin derivs. for diagnosis and therapy)

7440-54-2DP, Gadolinium, chelates with DOTA-biotin derivs., biological 10043-49-9DP, Gold 198, chelates with DOTA-biotin derivs., studies biological studies 10098-91-6DP, Yttrium 90, chelates with DOTA-biotin derivs., biological studies 13967-65-2DP, Holmium-166, chelates with DOTA-biotin derivs., biological studies 13968-53-1DP, Ruthenium 103, chelates with DOTA-biotin derivs., biological studies 13981-51-6DP, Mercury 197, chelates with DOTA-biotin derivs., biological studies 14119-09-6DP, Gallium 67, chelates with DOTA-biotin derivs., biological studies 14119-24-5DP, Osmium 191, chelates with DOTA-biotin derivs., biological studies 14133-76-7DP, Technetium 99, chelates with DOTA-biotin derivs., biological studies 14191-64-1DP, Praseodymium 142, chelates with DOTA-biotin derivs., biological studies 14265-75-9DP, Lutetium 177, chelates with DOTA-biotin derivs., biological studies 14265-85-1DP, Actinium 225, chelates with DOTA-biotin derivs., biological 14331-95-4DP, Ruthenium 105, chelates with DOTA-biotin derivs., biological studies 14378-26-8DP, Rhenium 188, chelates with DOTA-biotin derivs., biological studies 14391-11-8DP, Gold 199, chelates with DOTA-biotin derivs., biological studies 14391-19-6DP, Terbium 161, chelates with DOTA-biotin derivs., biological studies 14391-96-9DP, Scandium 47, chelates with DOTA-biotin derivs., biological studies 14687-25-3DP, Lead 203, chelates with DOTA-biotin derivs., biological studies 14885-78-0DP, Indium 113, chelates with DOTA-biotin derivs., biological studies 14913-49-6DP, Bismuth 212, chelates with DOTA-biotin derivs., biological studies 14913-89-4DP, chelates with DOTA-biotin derivs., biological studies 14914-68-2DP, Antimony 119, chelates with DOTA-biotin derivs., biological studies 14967-68-1DP, Palladium 103, chelates with DOTA-biotin derivs., biological studies 14981-64-7DP, Palladium 109, chelates with DOTA-biotin derivs., biological studies 14998-63-1DP, Rhenium 186, chelates with DOTA-biotin derivs., biological 15092-94-1DP, Lead 212, chelates with DOTA-biotin derivs., studies biological studies 15735-74-7DP, Platinum 197, chelates with DOTA-biotin derivs., biological studies 15750-15-9DP, Indium 111, chelates with DOTA-biotin derivs., biological studies 15756-62-4DP, Ruthenium 95, chelates with DOTA-biotin derivs., biological studies 15757-14-9DP, Gallium 68, chelates with DOTA-biotin derivs., biological studies 15757-86-5DP, Copper 67, chelates with DOTA-biotin derivs., biological studies 15758-35-7DP, Ruthenium 97, chelates with DOTA-biotin derivs., biological studies 15760-04-0DP, Silver 111, chelates with DOTA-biotin

derivs., biological studies 15765-78-3DP, Rhenium 189, chelates with DOTA-biotin derivs., biological studies 15766-00-4DP, Samarium 153, chelates with DOTA-biotin derivs., biological studies 294638-18-9P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(radionuclide conjugates containing DOTA-biotin derivs. for diagnosis and therapy)

T 170908-81-3P 192221-17-3P 192221-18-4P 192221-19-5P 245758-39-8P 294637-28-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(radionuclide conjugates containing DOTA-biotin derivs. for diagnosis and therapy)

IT 294638-18-9P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(radionuclide conjugates containing DOTA-biotin derivs. for diagnosis and therapy)

RN 294638-18-9 ZCAPLUS

The right of the result of the

IT 245758-39-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(radionuclide conjugates containing DOTA-biotin derivs. for diagnosis and therapy)

RN 245758-39-8 ZCAPLUS CN 1.4.7.10-Tetraazacyc

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[5-([3-48,48,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-coxopentyl]methylamino]ethyl]methylamino]-2-oxoethyl] (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

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THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 31 RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L79 ANSWER 17 OF 17 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:420994 ZCAPLUS Full-text

DOCUMENT NUMBER: 133:59099 TITLE: Preparation of vitronectin receptor antagonist

pharmaceuticals Harris, Thomas David; Rajodadhye, Milind

INVENTOR(S):

PATENT ASSIGNEE(S): Du Pont Pharmaceuticals Company, USA SOURCE: PCT Int. Appl., 300 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent. LANGUAGE: English FAMILY ACC. NUM. COUNT: 8

PATENT INFORMATION:

PA	TENT I	NO.			KINI)	DATE			APPL	ICAT:	ION I	NO.		D	ATE	
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WO	2000	0354	92		A2		2000	0622		WO 1	999-1	US30:	315		1	9991:	217
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                                                               20021126
PRIORITY APPLN. INFO.:
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                                                           P 19980331
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                                                            P 19981218
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                                          HS 1998-112831P
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                                                           A3 19990330
                                         US 1999-281209
                                                            A3 19990330
                                         WO 1999-US30315
                                                           W 19991217
OTHER SOURCE(S):
                       MARPAT 133:59099
AB
     Compds. (Q)d-Ln-Ch (Q is a residue having a quinolone-type moiety, d = 1-10,
     Ln is a linking group, Ch is a metal-bonding unit) were prepared for use in
     the diagnosis and treatment of cancer, methods of imaging tumors in a patient,
     and methods of treating cancer in a patient. The present invention also
     provides novel compds. useful for monitoring therapeutic angiogenesis
     treatment and destruction of new angiogenic vasculature. Thus, [3-[1-[3-[N-
     [3-[2-[N-(L-Asp-L-Asp)-3-
     aminopropoxy]ethoxy]ethoxy]propy1]carbamoy1]propanoylamino]propy1]-7-
     [(imidazol-2-vlamino)methyl]-4-oxo(3-hydroquinolyl)carbonylamino]-2- [[(2,4,6-
     trimethylphenyl)sulfonyl]amino]propanoic acid DOTA conjugate was prepared
     (claimed compound). Syntheses of radiopharmaceuticals, e.g.,
     99mTc(VnA) (tricine) (phosphine), where VnA represents the vitronectin receptor
     antagonist, are also described.
TC
    ICM A61K051-04
CC
    34-3 (Amino Acids, Peptides, and Proteins)
    Section cross-reference(s): 8, 27, 28, 63, 78
ΙT
    Angiogenesis
      Antitumor agents
    Atherosclerosis
    Radiopharmaceuticals
```

```
(preparation of vitronectin receptor antagonist pharmaceuticals)
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277316-58-2P
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
   (preparation of vitronectin receptor antagonist pharmaceuticals)
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TT 5704-04-1DP, Tricine, technetium-99 complexes 10098-91-6DP, complexes with vitronectin receptor binding conjugates, preparation 14133-76-7DP, complexes with vitronectin receptor binding conjugates, preparation 14265-75-9DP, complexes with vitronectin receptor binding conjugates, preparation 15750-15-9DP, complexes with vitronectin receptor binding conjugates, preparation 63995-70-0DP, TPPTS, technetium-99 complexes 277315-51-2P 277315-52-3P 277315-53-4P 277315-55-6P 277315-56-7P 277315-57-8P 277315-58-9P 277315-59-0P 277315-60-3P 277315-61-4P 277315-62-5P 277315-63-6P 277315-64-7P 277315-65-8P 277315-67-0P 277315-68-1P 277315-69-2P 277315-70-5P 277315-72-7P 277315-74-9P 277315-75-0P 277315-76-1P 277315-77-2P 277315-78-3P 277315-79-4P 277315-80-7P 277315-81-8DP, technetium-99 complexes 277316-60-6P 277316-61-7P 277316-62-8P 277316-63-9P 277316-64-0P 277316-65-1P 277316-66-2P 277316-67-3P 277316-68-4P 277316-69-5P 277316-71-9DP, technetium-99 complexes 277316-72-0DP, technetium-99 complexes 277316-73-1DP, technetium-99 complexes 277316-74-2DP, technetium-99 complexes 277316-75-3DP, technetium-99 complexes 277316-76-4DP, technetium-99 complexes 278172-91-1P 278172-92-2P 278172-93-3P 278172-94-4P 278172-95-5P 278172-96-6P 278172-97-7P 278172-98-8P 278172-99-9P 278173-00-5P 278173-01-6P 278173-02-7P 278173-03-8P 278173-04-9P 278173-05-0P 278173-06-1P 278173-07-2P 278173-08-3P 278173-09-4DP. gadolinium-labeled RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of vitronectin receptor antagonist pharmaceuticals)

TT 277316-47-9P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent) (preparation of vitronectin receptor antagonist pharmaceuticals)

277316-47-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[[(2S)-2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-vlamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2oxoethvll-, $\alpha, \alpha', \alpha''$ -tris(1,1-dimethylethyl) ester,

tris(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 277316-46-8 Absolute stereochemistry.

CMF C57 H85 N11 O13 S

PAGE 1-A

CM 2

CRN 76-05-1 CMF C2 H F3 O2

F_ _ CO21

CN

IT 277315-74-9P 277315-75-0P 2781/3-04-9P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of vitromectin receptor antagonist pharmaceuticals)

RN 277315-74-9 ZCAPLUS

 $\label{eq:continuity} 1, 4, 7, 10^{-} \text{Tetraazacyclododecane-1}, 4, 7^{-} \text{triacetic acid, } 10^{-} [2^{-} [3^{-} [3^{-} [([(2S)-2-\text{carboxy}-2^{-} [([(2,4,6-\text{trimethylphenyl}) \text{sulfonyl}] \text{amino}] \text{ethyl}] \text{amino}] \text{carbonyl}-7^{-} ([1H-\text{imidazol}-2-\text{ylamino}] \text{methyl}]-4-\text{oxo}-1(4H)-\text{quinolinyl}] \text{propyl}] \text{amino}]-2^{-}$

oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

RN 277315-75-0 ZCAPLUS

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[((2S)-CN 2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2oxoethyl]-, tris(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 277315-74-9 CMF C45 H61 N11 O13 S

PAGE 1-A

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 278173-04-9 ZCAPLUS

CN Yttrate(1-)-90%, [10-[2-[[3-[3-[1](2S)-2-carboxy-2-[[(2,4,6-trimethylphenyl)sulfonyl]mino]ethyl]mino]carbonyl]-7-[(1H-imidazol-2-ylmino)methyl]-4-oxo-1(4H)-quinollinyl]propyl]mino]-2-[oxo-k0]ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(4-)-kN1,kN4,kN7,kN10,kO1,kO4,kO7]-,hydrogen (9CI) (CA INDEX NAME)

PAGE 2-A

=> file registry FILE 'REGISTRY' ENTERED AT 10:26:43 ON 21 FEB 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 American Chemical Society (ACS)

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http://www.cas.org/support/stngen/stndoc/properties.html

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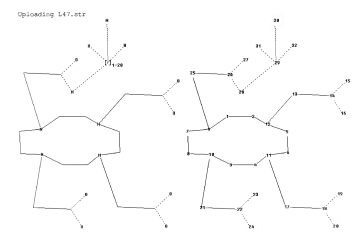
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 26:CLASS
 27:CLASS
 28:CLASS
 29:CLASS



chain nodes : 31 32 ring nodes :

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29-31 29-32 ring/chain bonds:
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Match level :

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 2:Atom
 3:Atom
 4:Atom
 5:Atom
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=> file zcaplus

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FILE COVERS 1907 - 21 Feb 2008 VOL 148 ISS 8 FILE LAST UPDATED: 20 Feb 2008 (20080220/ED)

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'OBI' IS DEFAULT SEARCH FIELD FOR 'ZCAPLUS' FILE

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
Structure attributes must be viewed using STN Express query preparation.
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          142 SEA FILE=REGISTRY ABB=ON PLU=ON L49 AND M/ELS
L50
L51
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          112 SEA FILE=REGISTRY ABB=ON PLU=ON L50 AND LNTH/PG
L56
L62
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             2 SEA FILE=ZCAPLUS ABB=ON PLU=ON (L51 OR L56) AND L62
L64
=> d stat que L65
1.25
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
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L50
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           203 SEA FILE=REGISTRY ABB=ON PLU=ON L49 NOT L50
L56
           112 SEA FILE=REGISTRY ABB=ON PLU=ON L50 AND LNTH/PG
L60
       641196 SEA FILE=ZCAPLUS ABB=ON PLU=ON ?TUMOUR?/BI OR ?TUMOR?/BI
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=> s L64-L65 not L79,L73,L74
L80
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=> d ibib abs hitind hitstr L80 1-32
L80 ANSWER 1 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2008:39770 ZCAPLUS Full-text
DOCUMENT NUMBER:
                        148:145033
TITLE:
                       Preparation of gastrin-releasing peptide compounds as
                       diagnostic imaging agents or radiotherapeutic agents
                       and methods of their use for treating prostate cancer
INVENTOR(S):
                        Cappelletti, Enrico; Lattuada, Luciano; Linder, Karen
                        E.; Marinelli, Edmund; Nanjappan, Palaniappa; Nunn,
                        Adrian D.; Raju, Natarajan; Ramalingam, Kondareddiar;
                        Swenson, Rolf E.; Tweedle, Michael; Maddalena, Marv
PATENT ASSIGNEE(S):
                       Bracco Imaging S.p.A., Italy
SOURCE:
                        U.S. Pat. Appl. Publ., 218pp., Cont.-in-part of U.S.
                        Ser. No. 352,156.
                        CODEN: USXXCO
DOCUMENT TYPE:
                        Patent.
LANGUAGE:
                       English
FAMILY ACC. NUM. COUNT: 6
PATENT INFORMATION:
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PA	PATENT NO.			KIN	D	DATE			APPLICATION NO.					DATE				
	US 2008008649			A1 20080110			US 2007-751337					20070521						
US	US 2004136906			A1 2004071			0715		US 2	003-	3415	77	20030113					
US	US 7226577			B2 20070605														
WO	WO 2004065407				A2 20040805			WO 2003-US41328				20031224						
WO	2004	0654	07		A3 20040923													
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US	2004	2532	25		A1		2004	1216		US 2	004-	8289	25		2	0040	420	
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AB The invention is related to novel gastrin-releasing peptide (GRP) compds. of formula M-N-O-P-G (M is an optical label or a metal chelator complexed with a radionuclide; N, P are null, an amino acid or other linking group; O is an amino acid; at least one of N, O, or P is a non- α -amino acid; G is a GRP receptor targeting peptide) which are useful as diagnostic imaging agents or radiotherapeutic agents. The invention is also related to methods for treating prostate tumors or of delaying the progression of prostate tumors, including, methods of treating bone or soft tissue metastases of prostate cancer, methods for treating hormone sensitive and hormone refractory prostate cancer, methods for delaying the progression of hormone sensitive prostate cancer, for facilitating combination therapy in patients with hormone sensitive prostate cancer and for decreasing aberrant vascular permeability in patients with hormone sensitive prostate cancer. Thus, DOTA-Gly-4-NHC6H4CO-L-Gln-L-Trp-L-Ala-L- Val-Gly-L-His-L-Leu-L-Met-NH2 (DOTA = 1,4,7,10tetraazacvclododecane- 1,4,7,10-tetraacetic acid residue) was prepared by the solid-phase method and complexed with 177Lu for cell binding, biodistribution and aberrant vascular permeability in LNCaP tumors studies.

INCL 424001690

CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 1, 8, 78

IT Antitumor agents

Combination chemotherapy

Human

Radiography

Radiotherapy

(preparation of gastrin-releasing peptide compds. for use as diagnostic imaging agents or radio therapeutic agents)

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                             721939-07-7P
                                                          721939-10-2P
721939-05-5P
                                           721939-08-8P
721939-11-3P
              721939-12-4P
                             721939-14-6P
                                           721939-16-8P
                                                          721939-17-9P
721939-18-0P
              721939-19-1P
                             721939-21-5P
                                           721939-23-7P
                                                          721939-25-9P
              721939-29-3P
721939-27-1P
                             721939-31-7P
                                           721939-33-9P
                                                          721939-35-1P
721939-37-3P
             722493-92-7P
                             722493-93-8P
                                           722493-94-9P
                                                          722493-95-0P
             722493-97-2P
722493-96-1P
                             722493-98-3P
                                           722493-99-4P
                                                          722494-00-0P
722494-01-1P
             722494-02-2P
                             808112-30-3P
                                          808112-31-4P
                                                          808112-32-5P
808112-33-6P
             808112-35-8P
                             808112-37-0P
                                          808112-39-2P
808112-41-6P
              808112-43-8P
                             808112-44-9P
                                          808112-45-0P
808112-46-1P
              808112-47-2P
                             808112-48-3P
                                           808112-49-4P
                                                          808112-50-7P
808112-51-8P
             808112-52-9P
                             808112-53-0P
                                           808112-54-1P
                                                          808112-55-2P
808112-56-3P
             808112-57-4P
                             808112-58-5P
                                           808112-59-6P
                                                          808112-60-9P
808112-61-0P
              808112-62-1P
                             808112-63-2P
                                           808112-64-3P
                                                          808112-65-4P
                                                          808112-71-2P
808112-67-6P
             808112-68-7P
                             808112-69-8P
                                           808112-70-1P
808112-72-3P
             808112-73-4P 803112-74-5P 808112-75-6P
808112-76-7P
              808113-15-7P
                            808113-16-8P 808113-17-9P
                                                          808113-18-0P
808113-19-1P
             808113-20-4P
                             808113-21-5P
                                                          808113-24-8P
                                           808113-23-7P
808113-25-9P
             808113-26-0P
                            808113-27-1P 808113-28-2P
                                                          808113-29-3P
809233-13-4P
             809233-16-7P
                            874367-58-5P 874534-72-2P
                                                          874534-73-3P
874537-63-0P
             913581-98-3P
                            913582-14-6P 913582-23-7P
                                                         913655-42-2P
913705-76-7P
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RL: DGN (Diagnostic use); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Blological study); PREP (Preparation); USES (Uses) (preparation of gastrin-releasing peptide compds. for use as diagnostic

IT 721937-82-2P 721937-90-2P 721937-92-4P

imaging agents or radio therapeutic agents)

808112-41-6P 808112-74-5P

RL: DGN (Diagnostic use); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of gastrin-releasing peptide compds. for use as diagnostic imaging agents or radio therapeutic agents)

RN 721937-82-2 ZCAPLUS

CN L-Methioninamide, N2-[[4-oxo-6-[4-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-1-piperazinyl]-3(H)-quinazolinyl]acetyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(9CI) (CA INDEX NAME)

PAGE 1-C

__SMe

RN 721937-90-2 ZCAPLUS

CN L-Methioninamide, N2-[[4-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-1-piperazinyl]acetyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl- (9CI) (CA INDEX NAME)

PAGE 1-B

PAGE 2-A

PAGE 2-B

RN 721937-92-4 ZCAPLUS

The thin oin a mide, N2-[[4-[2-[[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetrazacyclododec-l-yl]acetyl]amino]ethyl]-l-piperazinyl]acetyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(CA INDEX NAME)

PAGE 1-B

RN 808112-41-6 ZCAPLUS

CN L-Methioninamide, N2-[4-[[[2-[[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-l-yllacetyllamino]ethyllamino]carbonyllbenzoyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(GA INDEX NAME)

PAGE 1-B

→ SMe

PAGE 2-A

PAGE 2-B

RN 808112-74-5 ZCAPLUS

CN L-Methioninamide, N2-[4-[bis[2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]amino]ethyl]amino]-1,4-dioxobutyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(9CI)(CA INDEX NAME)

PAGE 1-B

 \sim SMe

HO₂C

PAGE 2-B

L80 ANSWER 2 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:1006173 ZCAPLUS Full-text

DOCUMENT NUMBER: 148:3337

TITLE: In vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromolecular MRI

contrast agent

Xu, Rongzuo; Wang, Yanli; Wang, Xuli; Jeong, Eun-Kee; AUTHOR(S): Parker, Dennis L.; Lu, Zheng-Rong

CORPORATE SOURCE: Dep. Parmaceutics and Pharmaceutical Chem., Univ.

Utah, Salt Lake City, UT, 84108, USA Experimental Biology and Medicine (Maywood, NJ, United

States) (2007), 232(8), 1081-1089

CODEN: EBMMBE; ISSN: 1535-3702

PUBLISHER: Society for Experimental Biology and Medicine

DOCUMENT TYPE: Journal

SOURCE:

AB

LANGUAGE: English

> Macromol. Gd(III) chelates are superior magnetic resonance imaging (MRI) contrast agents for blood pool and tamor imaging. However, their clin. development is limited by the safety concerns related to the slow excretion and long-term gadolinium tissue accumulation. A generation 6 PAMAM Gd(III) chelate conjugate with a cleavable disulfide spacer, PAMAM-G6-cystamine-(Gd-DO3A)1 was prepared as a biodegradable macromol. MRI contrast agent with rapid excretion from the body. T1 and T2 relaxivities of the contrast agent were 11.6 and 13.3 mM-1 sec-1 at 3T, resp. Blood pool and tumor contrast enhancement of the agent were evaluated in female nude mice bearing MDA-MB-231 human breast carcinoma xenograft swith a nondegradable conjugate PAMAM-G6-(Gd-DO3A) as a control. PAMAM-G6-cystamine-(Gd-DO3A) resulted in significant contrast enhancement in the blood for about 5 mins, and Gd-DO3A was released

from the conjugate and rapidly excreted via renal filtration after the disulfide spacer was cleaved. The nondegradable control had much longer blood circulation and excreted more slowly from the body. PAMAM-G6-cystamine-(Gd-D03A) also resulted in more prominent tumor contrast enhancement than the control. However, PAMAM-G6-cystamine-(Gd-D03A) demonstrated high toxicity due to the intrinsic toxicity of PAMAM dendrimers. In conclusion, although PAMAM-G6-cystamine-(Gd-D03A) showed some advantages compared with the nondegradable control. PAMAM dendrimers are not suitable carriers for biodegradable macromol. MRI contrast agents, due to their high toxicity.

CC 8-9 (Radiation Biochemistry) Section cross-reference(s): 9, 14

ST NRI contrast PANAM cystamine GdDO3A conjugate pharmacokinetics tumor imaging; mouse blood clearance MRI contrast agent disulfide spacer toxicity

IT Imaging

(NMR, tumor imaging using; in vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromol. MRI contrast agent)

IT 958259-88-6P

RL: BPN (Biosynthetic preparation); RCT (Reactant); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(in vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromol. MRI contrast agent)

T 99616-36-1P 150467-20-2P 958259-91-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(in vivo evaluation of a PAMAM-cystamine-(Gd-D03A) conjugate as a biodegradable macromol. MRI contrast agent)

IT 585531-76-6DP, PAMAM dendrimeric gadolinium complexes 958259-68-6DP, PAMAM dendrimeric gadolinium complexes

RL: SPN (Synthetic preparation); PREP (Preparation) (in vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromol. MRI contrast agent)

IT 958259-88-6P

RL: BPN (Biosynthetic preparation); RCT (Reactant); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(in vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromol. MRI contrast agent)

RN 958259-88-6 ZCAPLUS CN 1,4,7,10-Tetraazacyc

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-isothiocyanatoethyl)amino]-2-oxoethyl]- (CA INDEX NAME)

IT 150467-20-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(in vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromol. MRI contrast agent) $\,$

RN 150467-20-2 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2aminoethyl)amino]-2-oxoethyl]- (CA INDEX NAME)

II 958259-88-6UP, PAMAM dendrimeric gadolinium complexes RL: SPN (Synthetic preparation); PREP (Preparation)

(in vivo evaluation of a PAMAM-cystamine-(Gd-DO3A) conjugate as a biodegradable macromol. MRI contrast agent)

RN 958259-88-6 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-isothiocyanatoethyl)amino]-2-oxoethyl]- (CA INDEX NAME)

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 3 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:969732 ZCAPLUS Full-text

DOCUMENT NUMBER: 147:294732

TITLE: Polyamine-substituted ligands for use as contrast

agents

INVENTOR(S): Wolf, Markus; Bauder-Wust, Ulrike; Haberkorn, Uwe;

Eisenhut, Michael; Mier, Walter

PATENT ASSIGNEE(S): Germany SOURCE: U.S. Par

SOURCE: U.S. Pat. Appl. Publ., 20pp.
CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 2007202047 A1 20070830 US 2007-649503 20070104
PRIORITY APPLN. INFO:: US 2006-756352P P 20060105

OTHER SOURCE(S): MARPAT 147:294732

The present invention relates to a polyamine-substituted ligand for the preparation of a contrast agent derived from a chelating mol. selected from the group consisting of 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA) and diethylentriamine-pentaacetic acid (DTPA), wherein at least one of the carboxylic groups of the chelating mol. is reacted with an amine of formula HNR1R2 to form an amide bond, wherein R1, R2 are independently selected from the group consisting of H; (CH2)n-NR3R4; and R5; R3, R4 are independently selected from the group consisting of H; (CH2)m-NR6R7; and (CH2)m-1-CH3; R6, R7 are independently selected from the group consisting of H; and (CH2)o-1-CH3; n, m, o are independently 2, 3, or 4; R5 is of formula and optionally at least one of the carboxylic groups of the chelating mol. is further reacted with a monoalkylamine having 1 to 18 carbon atoms to form an amide bond; provided that at least one of R1, R2 is other than H. Furthermore, the invention relates to contrast agents for magnetic resonance imaging (MRI) comprising said ligands and in-vivo diagnostic methods based on MRI using said contrast agents.

INCL 424009363; 534015000; 540474000

CC 8-9 (Radiation Biochemistry)

ST polyamine substituted ligand gadolinium MRI tumor imaging

IT Imaging

(tumor; polyamine-substituted ligands for use as MRI contrast agents)

T 7440-54-2DP, Gadolinium, polyamine-substituted ligand conjugates 947391-67-5P 947391-68-6P 947391-69-7P 947391-70-0P 947391-71-1P 947391-72-2P 947391-73-3P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (polyamine-substituted liqands for use as MRI contrast agents)

IT 85503-20-4P 120131-72-8P 134935-60-7P 923952-46-9P 923952-47-0P 923952-48-1P 923952-49-2P 923952-50-5P 947337-79-3P 947337-80-6P 347337-81-7P 947337-82-9P 947337-83-9P 947337-84-0P 947337-85-1P 947337-86-2P 947337-87-3P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(polyamine-substituted ligands for use as MRI contrast agents)

IT 947331-70-0P 947391-71-1P 947391-72-2P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (polyamine-substituted ligands for use as MRI contrast agents)

RN 947391-70-0 ZCAPLUS

CN Gadolinium, [10-[2-[bis(2-aminoethyl)amino]-2-(oxo-kO)ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(8-)kN7, kN10,k01,k041- (CA INDEX NAME)

RN 947391-71-1 ZCAPLUS

CN Gadolinium, [10-[2-[(4-aminobuty1)(3-aminopropy1)amino]-2-(oxo-KO)ethy1]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)-KN1,KN4,KN7,KN10,KO1,KO4,KO7]-(CA INDEX NAME)

RN 947391-72-2 ZCAPLUS

CN Gadolinium, [10-[2-[(3-aminopropyl)[4-[(3-aminopropyl)amino]butyl]amino]-2-(oxo-KO)ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)-KN1,KN4,KN7,KN10,KO1,KO4,KO7]-(CA INDEX NAME)

- IT 947337-81-7P 947337-82-8P 947337-83-9P 947337-86-2P 947337-87-3P
 - RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
- (polyamine-substituted ligands for use as MRI contrast agents)
- RN 947337-81-7 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[bis(2aminoethyl)amino]-2-oxoethyl]- (CA INDEX NAME)

- RN 947337-82-8 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-aminobuty1)(3-aminopropy1)amino]-2-oxoethyl]- (CA INDEX NAME)

- RN 947337-83-9 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(3-aminopropyl)[4-[(3-aminopropyl)amino]butyl]amino]-2-oxoethyl]- (CA INDEX NAME)

- RN 947337-86-2 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-aminobuty1) (3-aminopropy1)amino]-2-oxoethyl]-, 1,4,7-tris(1,1-dimethylethyl) ester (CA INDEX NAME)

$$\begin{array}{c} \text{CH2} & \text{CH2} & \text{3} = \text{HH2} \\ \text{CH2} & \text{CH2} & \text{CH2} & \text{4} = \text{HH2} \\ \text{CH2} & \text{CH2} & \text{CH2} & \text{4} = \text{HH2} \\ \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} & \text{4} = \text{HH2} \\ \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} \\ \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} \\ \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} & \text{CH2} \\ \end{array}$$

RN 947337-87-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(3aminopropyl) [4-[(3-aminopropyl)amino]butyl]amino]-2-oxoethyl]-, 1,4,7-tris(1,1-dimethylethyl) ester (CA INDEX NAME)

L80 ANSWER 4 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:960536 ZCAPLUS Full-text DOCUMENT NUMBER: 147:464287

TITLE: Polymer-based elemental tags for sensitive bioassays AUTHOR(S): Lou, Xudong; Zhang, Guohua; Herrera, Isaac; Kinach,

Robert; Olga, Ornatsky; Baranov, Vladimir; Nitz, Mark;

Winnik, Mitchell A.

CORPORATE SOURCE: Institute of Biomaterials and Biomedical Engineering, University of Toronto, Toronto, ON, M5S 3G9, Can.

Angewandte Chemie, International Edition (2007),

46(32), 6111-6114, \$6111/1-\$6111/5 CODEN: ACIEF5; ISSN: 1433-7851

Wiley-VCH Verlag GmbH & Co. KGaA

DOCUMENT TYPE: Journal

LANGUAGE: English

A water-soluble polymer bearing multiple metal-chelating ligands has been used as a tag for bioassays with inductively coupled plasma mass spectrometry. The tag was covalently conjugated to antibodies, and the polymer-antibody constructs were loaded with lanthanide ions (Ln3+) and used for the simultaneous assay of five orthogonally labeled antibodies against cell surface antigens that differ in abundance by more than two orders of magnitude.

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 14, 15, 35

IT Acute monocytic leukemia Acute myeloid leukemia Chelating agents Chelation

SOURCE:

PUBLISHER:

Diagnostic agents

Human

Immunoassay

Molecular recognition

Protein-protein interaction

Tumor markers

(water-soluble polymer bearing multiple metal-chelating ligands tag for bioassays with inductively coupled plasma mass spectrometry applied to leukemia)

IT 115597-84-7 150463-52-8D, t-Bu, dithiobenzoate terminated

173308-19-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(water-soluble polymer bearing multiple metal-chelating ligands tag for bioassays with inductively coupled plasma mass spectrometry applied to leukemia)

IT 100-46-9DP, Benzenemethanamine, reaction with acrylamide/acrylic acid polymer, preparation 150467-20-2DP, reaction with

acrylamide/acrylic acid polymer 173308-19-5DP, reaction with

acrylamide/acrylic acid polymer

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(water-soluble polymer bearing multiple metal-chelating ligands tag for bioassays with inductively coupled plasma mass spectrometry applied to leukemia)

IT 173308-19-5

RL: RCT (Reactant); RACT (Reactant or reagent)

(water-soluble polymer bearing multiple metal-chelating ligands tag for bloassays with inductively coupled plasma mass spectrometry applied to leukemia)

RN 173308-19-5 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]-, 1,4,7-tris(1,1-dimethylethyl) ester (CA INDEX NAME)

$$\begin{array}{c} \text{CH}_2 - \text{C} \\ \text{CH}_2 - \text{C} \\ \text{OB} \\ \text{CH}_2 - \text{C} \\ \text{OB} \\ \text{OB$$

IT 150467-20-20P, reaction with acrylamide/acrylic acid polymer 173309-19-50P, reaction with acrylamide/acrylic acid polymer

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(water-soluble polymer bearing multiple metal-chelating ligands tag for bioassays with inductively coupled plasma mass spectrometry applied to leukemia)

RN 150467-20-2 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2aminoethyl)amino]-2-oxoethyl]- (CA INDEX NAME)

RN 173308-19-5 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]-, 1,4,7-tris(1,1-dimethylethyl) ester (CA INDEX NAME)

$$\begin{array}{c} \text{t-Buo-} \overset{\circ}{\text{U}} = \text{CH}_2 - \overset{\circ}{\text{U}} = \text{OBu-t} \\ & \text{CH}_2 - \overset{\circ}{\text{U}} = \text{OBu-t} \\ & \text{CH}_2 - \overset{\circ}{\text{U}} = \text{OBu-t} \\ & \text{CH}_2 - \text{CH}_2 - \text{CH}_2 - \text{NH}_2 \\ \end{array}$$

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 5 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:545418 ZCAPLUS Full-text

DOCUMENT NUMBER: 147:206685

TITLE: Noninvasive Visualization of Pharmacokinetics,

Biodistribution and Tumor Targeting of Poly[N-(2-hydroxypropyl)methacrylamide] in Mice Using

Contrast Enhanced MRI

AUTHOR(S): Wang, Yanli; Ye, Furong; Jeong, Eun-Kee; Sun, Yongen;

Parker, Dennis L.; Lu, Zheng-Rong

CORPORATE SOURCE: Department of Pharmaceutics and Pharmaceutical

Chemistry, University of Utah, Salt Lake City, UT,

84108, USA

SOURCE: Pharmaceutical Research (2007), 24(6), 1208-1216

CODEN: PHREEB; ISSN: 0724-8741

PUBLISHER: Springer DOCUMENT TYPE: Journal

LANGUAGE: Journal English

AB Purpose: To study a non-invasive method of using contrast enhanced magnetic resonance imaging (MRI) to visualize the real-time pharmacokinetics, biodistribution and tumor accumulation of paramagnetically labeled poly[N-(2-hydroxypropyl)methacrylamide] (PHPMA) copolymer conjugates with different mol. wts. and spacers in tumor-bearing mice. Materials and Methods: Paramagnetically labeled HPMA copolymer conjugates were synthesized by free radical copolymn. of HPMA with monomers containing a chelating licend.

radical copolymm. or HPAN with monomers containing a chelating ligand, followed by complexation with Gd(OAc)3. A stable paramagnetic chelate, Gd-DO3A, was conjugated to the copolymers via a degradable spacer GlyPheLeuGly

and a non-degradable spacer GlyGly, resp. The conjugates with mol. wts. of 28, 60 and 121 kDa and narrow mol. weight distributions were prepared by fractionation with size exclusion chromatog. The conjugates were injected into athymic nude mice bearing MDA-MB-231 human breast carcinoma xenografts via a tail vein. MR images were acquired before and at various time points after the injection with a 3D FLASH sequence and a 2D spin-echo sequence at 3T. Pharmacokinetics, biodistribution and tumor accumulation of the conjugates were visualized based on the contrast enhancement in the blood, major organs and tumor tissue at various time points. The size effect of the conjugates was analyzed among the conjugates. Results: Contrast enhanced MRI resulted in a real-time, three-dimensional visualization of blood circulation, pharmacokinetics, biodistribution and tumer accumulation of the conjugates, and the size effect on these pharmaceutical properties. HPMA copolymer conjugates with high mol, weight had a prolonged blood circulation time and high passive tumor targeting efficiency. Non-biodegradable HPMA copolymers with mol. wts. higher than the threshold of renal filtration demonstrated higher efficiency for tumpr drug delivery than biodegradable poly(L-glutamic acid). Conclusions: Contrast enhanced MRI is an effective method for noninvasive visualization of in vivo properties of the paramagnetically labeled polymer conjugates in preclin. studies.

- CC 8-9 (Radiation Biochemistry)
- ST contrast MRI gadolinium hydroxypropyl methacrylamide copolymer pharmacokinetics tumor imaging
- IT Human
 - Pharmacokinetics

(MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT Imaging agents

(NMR contrast; MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT Imaging

(NMR; MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT Mammary gland, neoplasm

(carcinoma; MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT Carcinoma

(mammary; MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT Imaging

(tumor; MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT 944834-63-3P 944834-65-5P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

- IT 21442-01-3, N-(2-Hydroxypropyl)methacrylamide 57950-79-5 100424-71-3 912576-20-6 944731-76-4
 - RL: RCT (Reactant); RACT (Reactant or reagent)

(MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tomor targeting)

- 944731-74-2P 944731-75-3P
- RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting)

IT 944834-63-3P 944834-65-5P

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tomor targeting)

RN 944834-63-3 ZCAPLUS

CN Gadolinium, [N-(2-methyl-1-oxo-2-propen-1-yl)glycyl-N-[6-[[2-[4,7,10-tris[(Carboxy-KO)methyl]-1,4,7,10-tetraazacyclododec-1-yl-KN1,KN4,KN7,KN10]acetyl-

κO]amino]hexyl]glycinamidato(3-)]-, polymer with

N-(2-hydroxypropy1)-2-methy1-2-propenamide (CA INDEX NAME)

CM 1

CRN 944834-62-2 CMF C30 H49 Gd N8 O10

CCI CCS

PAGE 1-A

PAGE 1-B

CM 2

CRN 21442-01-3 CMF C7 H13 N O2

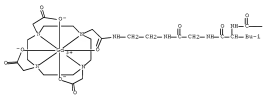
RN 944834-65-5 ZCAPLUS
CN Gadolinium, [N-(2-methyl-1-oxo-2-propen-1-yl)glycyl-L-phenylalanyl-L-leucyl-N-[2-[12-[4,7,10-tris]((carboxy-KO)methyl]-1,4,7,10-tetrazacyclododec-1-yl-KNl,KNA,KN7,KNl0]acetyl-KO]amino|ethyl]glycinamidato(3-)]-, polymer with
N-(2-hydroxypropyl)-2-methyl-2-propenamide (CA INDEX NAME)

CM 1

CRN 944834-64-4 CMF C41 H61 Gd N10 O12

CCI CCS

PAGE 1-A



PAGE 1-B

CM 2

CRN 21442-01-3 CMF C7 H13 N O2

CN

RL: RCT (Reactant); RACT (Reactant or reagent) (MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics and tumor targeting) RN 912576-20-6 ZCAPLUS 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6aminohexyl)amino]-2-oxoethyl]-, 2,2,2-trifluoroacetate (1:1) (CA INDEX NAME) CM 1 CRN 889140-15-2 CMF C22 H42 N6 O7

CM 2 CRN 76-05-1 CMF C2 H F3 O2

RN 944731-76-4 ZCAPLUS

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2-CN aminoethyl)amino]-2-oxoethyl]-, 2,2,2-trifluoroacetate (1:1) (CA INDEX NAME)

CM 1

CRN 150467-20-2 CMF C18 H34 N6 O7

CM 2

CRN 76-05-1 CMF C2 H F3 O2

IT 944731-74-2P 944731-75-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (MRI visualization of polyhydroxypropyl methacrylamide pharmacokinetics

and tempor targeting)

RN 944731-74-2 ZCAPLUS CN Glycinamide, N-(2-met

Glycinamide, N-(2-methyl-1-oxo-2-propen-1-yl)glycyl-N-[6-[[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]amino]hexyl]-(CA INDEX NAME)

PAGE 1-B

RN 944731-75-3 ZCAPLUS

CN Glycinamide, N-(2-methyl-1-oxo-2-propen-1-yl)qlycyl-L-phenylalanyl-Lleucyl-N-[2-[[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1yl]acetyl]amino]ethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 27 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 6 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 2007:402215 ZCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 146:421772

TITLE: Biotin diamino derivatives and their conjugates with

macrocyclic chelating agents

INVENTOR(S): Carminati, Paolo; Ginanneschi, Mauro; Paganelli,

Giovanni; Chinol, Marco

PATENT ASSIGNEE(S): Sigma-Tau Industrie Farmaceutiche Riunite S.p.A.,

Italy

PCT Int. Appl., 25pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Pat.ent. LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.		KIND	DATE	APPLI		DATE						
WO 200703943	37	A1	20070412	WO 20		20060918						
W: AE,	AG, AL,	AM, AT	AU, AZ,	BA, BB,	BG, BR, BW,	BY, BZ	, CA, CH,					
CN,	CO, CR,	CU, C2	, DE, DK,	DM, DZ,	EC, EE, EG,	ES, FI	, GB, GD,					
GE,	GH, GM,	HN, HE	R, HU, ID,	IL, IN,	IS, JP, KE,	KG, KM	, KN, KP,					
KR,	KZ, LA,	LC, LE	, LR, LS,	LT, LU,	LV, LY, MA,	MD, MG	, MK, MN,					
MW,	MX, MY,	MZ, NA	, NG, NI,	NO, NZ,	OM, PG, PH,	PL, PT	, RO, RS,					

GI

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RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW, ZW

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CT, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GM, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TU, TM

PRIORITY APPLM. INFO::

OTHER SOURCE(S):

CASREACT 146:421772; MARPAT 146:421772
```

HN H NHR' T

$$\Gamma = -CO - W$$

$$V$$

$$V$$

$$V$$

$$V$$

$$V$$

$$V$$

$$V$$

$$V$$

$$V$$

Biotin diamino derivs. I [A = CH2, C:O; B = H, CHO, CO2H; C = (CH2)c; D = AB (CH2)d; W = C1-12-alkylene, C2-12-alkenylene, functionalized polyethylene glycol, C6-10-aromatic residue, glucofuranosyl residue; R = linear or branched C1-4-alkyl, cycloalkyl, heterocycle, (CH2)qT; T = SMe, OH, CO2H; Q = 0, 1, 2; R', R'' = L; M = (CH2)m; P = (CH2)p; X = H, CH2U, (CHJ)oZ; Y = H, (un)branched C1-44-alkyl, (CH2)mCO2H; U = Me, Et, C6H4NH2-4; Z = NH2, NHC(:NH)NH2, SR2, 5or 6-membered heterocycle containing one or more O, S, NR1; R1 = H, linear or branched C1-4-alkyl; R2 = linear or branched C1-4-alkyl; J = H, Me, Et; n = 4 -12; a, b = 0 - n-1; c, d = 3 - 10; m = 1 - 3; o = 1 - 5; p = 2, 3] are described. Processes for their preparation, and their uses for the preparation of conjugates with radionuclides for use in human and animal therapy and diagnostics, particularly for the diagnosis and therapy of pathol. conditions such as tumors. Thus, I [A = W = CH2, B = R = H, Q = (CH2)6, c = d = 3, R' = R'' = 4,7,10-tri(carboxymethyl) - 1,4,7,10-tetrazacyclododecane-1acetyl] was prepared from reduced biotin N-hexylamide via acylation with N,Nbis[3-[(9- fluorenvlmethoxycarbonvl)amino]propvl]glycine potassium sulfate, deprotection with piperidine in DMF and acylation with DOTA.

CC 26-8 (Biomolecules and Their Synthetic Analogs) Section cross-reference(s): 8, 63, 78

IT Radiopharmaceuticals

(antitumor; preparation of biotin conjugates with macrocyclic amines for therapeutic use as chelating agents)

IT Ansitumor agents

Neoplasm

(radiopharmaceuticals; preparation of biotin conjugates with macrocyclic amines for therapeutic use as chelating agents)

IT 934166-99-1P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological

RN

study); PREP (Preparation); USES (Uses)

(preparation of biotin conjugates with macrocyclic amines for therapeutic use as chelating agents)

IT 934166-99-1P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of biotin conjugates with macrocyclic amines for therapeutic use as chelating agents)

934166-99-1 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10,10'-[[[2-[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazo1-4-y1]pentyl]amino]hexyl]amino]-2-oxoethyl]imino]bis[3,1-propanediylimino(2-oxo-2,1-ethanediyl)]]bis (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 2-A

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 7 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:377649 ZCAPLUS Full-text

DOCUMENT NUMBER: 147:66371

TITLE: Physicochemical and MRI characterization of

Gd3+-loaded polyamidoamine and hyperbranched

dendrimers

AUTHOR(S): Jaszberenyi, Zoltan; Moriggi, Loieck; Schmidt,
Philipp; Weidensteiner, Claudia; Kneuer, Rainer;

Merbach, Andre E.; Helm, Lothar; Toth, Eva

CORPORATE SOURCE: Institut des Sciences et Ingenierie Chimiques, Ecole

Polytechnique Federale de Lausanne, ISIC, BCH,

Lausanne, 1015, Switz.

SOURCE: JBIC, Journal of Biological Inorganic Chemistry

(2007), 12(3), 406-420 CODEN: JJBCFA: ISSN: 0949-8257

Springer GmbH

DOCUMENT TYPE: Journal

PUBLISHER:

LANGUAGE: English

Generation 4 polyamidoamine (PAMAM) and, for the first time, hyperbranched poly(ethylene imine) or polyglycerol dendrimers have been loaded with Gd3+ chelates, and the macromol, adducts have been studied in vitro and in vivo with regard to MRI contrast agent applications. The Gd3+ chelator was either a tetraazatetracarboxylate DOTA-pBn4- or a tetraazatricarboxylate monoamide DO3A-MA3- unit. The water exchange rate was determined from a 170 NMR and 1H Nuclear Magnetic Relaxation Dispersion study for the corresponding monomer analogs [Gd(DO3A-AEM)(H2O)] and [Gd(DOTA-pBn-NH2)(H2O)]- (k = 3.4 and $6.6 \times$ 106 s-1, resp.), where H3DO3A-AEM is {4-[(2-acetylaminoethylcarbamoyl)methyl]-7,10-bis(carboxymethyl-1,4,7,10- tetraazacyclododec-1-yl)}-acetic acid and H4DOTA-pBn-NH2 is 2-(4-aminobenzyl)-1,4,7,10-tetraazacyclododecane-1,4,7,10tetraacetic acid. For the macromol. complexes, variable-field proton relaxivities have been measured and analyzed in terms of local and global motional dynamics by using the Lipari-Szabo approach. At frequencies below 100 MHz, the proton relaxivities are twice as high for the dendrimers loaded with the neg. charged Gd(DOTA-pBn) - in comparison with the analogous mol. bearing the neutral Gd(DO3A-MA). We explained this difference by the different rotational dynamics: the much slower motion of Gd(DOTA-pBn) -- loaded dendrimers is likely related to the neg, charge of the chelate which creates more rigidity and increases the overall size of the macromol. compared with dendrimers loaded with the neutral Gd(DO3A-MA). Attachment of poly(ethylene glycol) chains to the dendrimers does not influence relaxivity. Both hyperbranched structures were found to be as good scaffolds as regular PAMAM dendrimers in terms of the proton relaxivity of the Gd3+ complexes. The in vivo MRI studies on tumor-bearing mice at 4.7 T proved that all dendrimeric

complexes are suitable for angiog. and for the study of vasculature parameters like blood volume and permeability of tumor vessels.

CC 6-7 (General Biochemistry)

Section cross-reference(s): 1, 63

T 941280-58-6P

RL: DGN (Diagnostic use); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (nod c,att o d,mov,joi; physicochem, and MRI characterization of

(nod c,att o d,mov,joi; physicochem. and MRI characterization of Gd3+-loaded polyamidoamine and hyperbranched dendrimers)

Gd3+-loaded polyamidoamine and hyperbranched dendrimers)
T 9002-98-6DP, reaction products with PAMAM, gadolinium complexes

9004-74-40P, PAMAM-PEI derivs. 25618-55-7DP, Polyglycerol, amine-functionalized 26937-01-9DP, reaction products with polyethylenimine, gadolinium complexes 120041-09-0DP, PAMAM-PEI gadolinium dendritic derivs. 123317-52-2DP, PAMAM-PEI gadolinium ethoxylated/polyglycerol dendritic derivs. 940951-69-3P

941280-59-7P

RL: DGN (Diagnostic use); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (physicochem. and MRI characterization of Gd3+-loaded polyamidoamine

and hyperbranched dendrimers)

IT 941280-58-6P

RL: DGN (Diagnostic use); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(nod c,att o d,mov,joi; physicochem. and MRI characterization of Gd3+-loaded polyamidoamine and hyperbranched dendrimers)

RN 941280-58-6 ZCAPLUS

CN Gadolinium, [10-[2-[[2-(acetylamino)ethyl]amino]-2-(oxo-KO)ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)-KNI,KN4,KN7,KN10,KO1,KO4,KN7]aqu

a- (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

IT 940961-69-3P

RL: DGN (Diagnostic use); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(physicochem. and MRI characterization of Gd3+-loaded polyamidoamine and hyperbranched dendrimers)

RN 940961-69-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-(acetvlamino)ethvl]amino]-2-oxoethvl]- (CA INDEX NAME)

REFERENCE COUNT:

44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 8 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:249358 ZCAPLUS Full-text DOCUMENT NUMBER: 146:501325

SOURCE:

TITLE: Synthesis of DOTA-conjugated multivalent cyclic-RGD

peptide dendrimers via 1,3-dipolar cycloaddition and their biological evaluation: implications for tumor

targeting and tumor imaging purposes

AUTHOR(S): Dijkgraaf, Ingrid; Rijnders, Anneloes Y.; Soede,
Annemieke; Dechesne, Annemarie C.; Van Esse, G. Wilma;

Brouwer, Arwin J.; Corstens, Frans H. M.; Boerman, Otto C.; Rijkers, Dirk T. S.; Liskamp, Rob M. J.

CORPORATE SOURCE: Department of Medicinal Chemistry and Chemical
Biology, Utrecht Institute for Pharmaceutical

Sciences, Utrecht University, Utrecht, 3508 TB, Neth.
Organic & Biomolecular Chemistry (2007), 5(6), 935-944

CODEN: OBCRAK; ISSN: 1477-0520

PUBLISHER: Royal Society of Chemistry

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 146:501325

AB The design and synthesis of a series of $\alpha V\beta 3$ integrin-directed monomeric, dimeric and tetrameric cyclo[Arg-Glv-Asp-d-Phe-Lvs] dendrimers using "click chemical" is described. It was found that the unprotected N-.vepsiln.-azido derivative of cyclo[Arg-Gly-Asp-d-Phe-Lys] underwent a highly chemoselective conjugation to amino acid-based dendrimers bearing terminal alkynes using a microwave-assisted Cu(i)-catalyzed 1,3-dipolar cycloaddn. The $\alpha V\beta 3$ binding characteristics of the dendrimers were determined in vitro and their in vivo $\alpha V \beta 3$ targeting properties were assessed in nude mice with s.c. growing human SK-RC-52 tumors. The multivalent RGD-dendrimers were found to have enhanced affinity toward the $\alpha V\beta 3$ integrin receptor as compared to the monomeric derivative as determined in an in vitro binding assay. In case of the DOTAconjugated 111In-labeled RGD-dendrimers, it was found that the radiolabeled multimeric dendrimers showed specifically enhanced uptake in $\alpha V\beta 3$ integrin expressing tumors in vivo. These studies showed that the tetrameric RGDdendrimer had better tumor targeting properties than its dimeric and monomeric congeners.

CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 1, 28

ST DOTA cyclic RGD peptide conjugated dendrimer prepn tumor imaging; cyclic RGD peptide solid phase prepn DOTA dipolar cycloaddn

IT Cvcloaddition reaction

(1,3-dipolar; preparation and tumor targeting and imaging use of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using microwave-assisted dipolar cycloaddn. as the key step for the conjugation)

IT Microwave

(irradiation; preparation and tumor targeting and imaging use of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using microwave-assisted dipolar cycloaddn. as the key step for the conjugation)

IT Antitumor agents

Human

Pharmacokinetics

(preparation and tumor targeting and imaging use of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using

microwave-assisted dipolar cycloaddn. as the key step for the conjugation)

IT RGD peptides

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and tumor targeting and imaging use of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using

IT

ΤТ

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microwave-assisted dipolar cycloaddn, as the key step for the
   conjugation)
Imaging
Imaging agents
   (tumex; preparation and tumex targeting and imaging use
   of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using
   microwave-assisted dipolar cycloaddn, as the key step for the
   conjugation)
Integrins
RL: BSU (Biological study, unclassified); BIOL (Biological study)
   (ανβ3; preparation and tumer targeting and imaging use
   of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using
   microwave-assisted dipolar cycloaddn, as the key step for the
   conjugation)
936125-37-0P 936125-39-2P 936235-89-1P
RL: DGN (Diagnostic use); PAC (Pharmacological activity); PKT
(Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study);
PREP (Preparation); USES (Uses)
   (preparation and tumor targeting and imaging use of
   DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using
   microwave-assisted dipolar cycloaddn, as the key step for the
   conjugation)
99-06-9, 3-Hydroxybenzoic acid, reactions 99-10-5, 3,5-Dihydroxybenzoic
acid 106-96-7, Propargyl bromide 107-15-3, 1,2-Ethanediamine, reactions 29022-11-5, Fmoc-Gly-OH 39684-80-5, tert-Butyl
(2-bromoethyl)carbamate 71989-14-5 71989-26-9 86123-10-6
137076-54-1 154445-77-9
RL: RCT (Reactant); RACT (Reactant or reagent)
   (preparation and tumor targeting and imaging use of
   DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using
   microwave-assisted dipolar cycloaddn. as the key step for the
   conjugation)
2150-44-9P, 3,5-Dihydroxybenzoic acid, methyl ester 19438-10-9P,
3-Hydroxybenzoic acid, methyl ester 57260-73-8P 85607-73-4P
160893-68-5P 184916-28-7P 250612-44-3P 664334-21-8P 680572-35-4P
768387-51-5P 866088-22-4P 936125-14-3P 936125-18-7P 936125-20-1P
936125-22-3P 936125-24-5P 936125-26-7P
936125-28-9P 936125-31-4P 942131-93-3P 942131-95-5P 942131-99-9P 942132-29-8P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
   (preparation and tumos targeting and imaging use of
   DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using
   microwave-assisted dipolar cycloaddn. as the key step for the
   conjugation)
868845-24-3P
              868845-25-4P 936125-33-6P
RL: SPN (Synthetic preparation); PREP (Preparation)
   (preparation and tumor targeting and imaging use of
   DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using
   microwave-assisted dipolar cycloaddn. as the key step for the
   conjugation)
936125-37-0P 936125-39-2P
```

RL: DGN (Diagnostic use); PAC (Pharmacological activity); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and tumor targeting and imaging use of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using microwave-assisted dipolar cycloaddn. as the key step for the conjugation)

RN 936125-37-0 ZCAPLUS

CN Cyclo[L-arginylglycyl-L-\(\alpha\)-aspartyl-D-phenylalanyl-6-[4-[[3-[[2-[[2-[4,7,10-tris(carboxymethyl)-1,4,7,10-tetrazacyclododec-l-yl]acetyl]amino]ethyl]amino]carbonyl]phenoxy]methyl]-H-1,2,3-triazol-l-yl]-L-norleucyl] (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

- RN 936125-39-2 ZCAPLUS
- CN Cyclo(L-arginylglycyl-L-a-aspartyl-D-phenylalanyl-L-norleucyl),
 56,5'-6[[5-[[2-[q-14,7].o-tris(carboxymethyl)-1,4,7,10tetraazacyclododec-1-yl]acetyl]amino]ethyl]amino]carbonyl]-1,3phenylene]bis(oxymethylene-1H-1,2,3-triazole-4,1-diyl)]bis(CA INDEX
 NAME)

PAGE 3-A

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___(CH2)4

PAGE 3-B

IT 936125-22-3P 936125-24-5P 936125-28-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and tumor targeting and imaging use of DOTA-conjugated multivalent cyclic-RGD peptide dendrimers using microwave-assisted dipolar cycloaddn. as the key step for the conjugation)

RN 936125-22-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-oxo-2-[[2-[3-(2-propn-1-yloxy)benzoyl]amino]ethyl]amino]ethyl]-, 1,4,7-tris(1,1-dimethylethyl) ester (CA INDEX NAME)

RN 936125-24-5 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[3,5-bis(2propyn-1-yloxy)benzoyl]amino]ethyl]amino]-2-oxoethyl]--, 1,4,7-trie(1,1-dimethylethyl) ester (CA INDEX NAME)

RN 936125-28-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[3,5-bis[2-[3,5-bis[2-propyn-1-yloxy)]benzoyl]amino]ethoxy]benzoyl]amino]ethoyl]benzoylamino]ethoyl]-1,4,7-tris[1,1-dimethylethyl] ester (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 2-A

REFERENCE COUNT:

THERE ARE 80 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 9 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:230231 ZCAPLUS <u>Full-text</u>

80

DOCUMENT NUMBER: 146:288424
TITLE: Non-invasi

TITLE: Non-invasive diagnostic agents of cancer and methods of diagnosing cancer, especially leukemia and lymphoma INVENTOR(S): Norenberg, Jeffrey P. PATENT ASSIGNEE(S): USA

PATENT ASSIGNEE(S): USA SOURCE: U.S. Pat. Appl. Publ., 19pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. US 2007048216 A1 20070301 US 2006-507846 20060822 US 2005-710665P P 20050823 PRIORITY APPLN. INFO.:

AB The present invention is directed to novel non-invasive diagnostic tools to image cancers, especially, leukemia and non-Hodgkin's lymphomas (NHL) with minimal toxicity in vivo. The present invention represents a clear advance in the art which presently relies on tissue biopsy for diagnoses of these cancers. The novel imaging probe is capable of detecting precancerous cells, as well as their metastatic spread in tissues. This represents a quantum step forward in the diagnosis and staging of NHL using non-invasively mol. imaging techniques. This novel probe will also be useful to monitor patients response to chemotherapy treatments and other interventions or therapies used in the treatment of NHL. Compds. according to the present invention may be used as diagnostic tools for a number of conditions and diseases states as well as therapeutic agents for treating such conditions and disease states.

INCL 424001110; 534011000; 534014000

1-6 (Pharmacology)

Section cross-reference(s): 4, 8, 63

IT Acute lymphocytic leukemia Acute myeloid leukemia

Acute promyelocytic leukemia Adult T-cell leukemia

Anti-inflammatory agents Anti-ischemic agents

Antidiabetic agents Antirheumatic agents

Antitumor agents

Arthritis Autoimmune disease

Blood analysis

Cardiopulmonary bypass

Diabetes mellitus Diagnostic agents

Drug toxicity

Hairy cell leukemia

Hematopoiesis

Human Imaging

Immunity

Inflammation

Inflammatory bowel diseases

Ischemia

Monocytic leukemia Multiple sclerosis

Mveloid leukemia

Myocardial infarction

Neoplasm

Osteoarthritis

Polymorphonuclear leukocyte

Psoriasis

Respiratory distress syndrome Rheumatoid arthritis

Skin, disease

Stem cell Transplant rejection Uveitis

Wart.

(non-invasive diagnostic agents of cancer and methods of diagnosing cancer, especially leukemia and lymphoma)

927833-57-6 927833-59-8

> RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); DGN (Diagnostic use); PKT (Pharmacokinetics); BIOL (Biological study); USES (Uses)

(non-invasive diagnostic agents of cancer and methods of diagnosing cancer, especially leukemia and lymphoma)

927833-57-6

RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); DGN (Diagnostic use); PKT (Pharmacokinetics); BIOL (Biological study); USES (Uses)

(non-invasive diagnostic agents of cancer and methods of diagnosing cancer, especially leukemia and lymphoma)

927833-57-6 ZCAPLUS RN

CN Lutetium-177Lu, [10-[2-[[4-[5-[(4-bromopheny1)methy1]-3-(3,5dichlorophenyl)-5-methyl-2,4-dioxo-1-imidazolidinyl]butyl]amino]-2-(oxoκO)ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)κN1,κN4,κN7,κN10]- (CA INDEX NAME)

L80 ANSWER 10 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2007:78033 ZCAPLUS Full-text

DOCUMENT NUMBER: 147:517229

TITLE: PET imaging of apoptosis with 64Cu-labeled

streptavidin following pretargeting of

phosphatidylserine with biotinylated annexin-V AUTHOR(S): Cauchon, Nicole; Langlois, Rejean; Rousseau, Jacques

A.; Tessier, Guillaume; Cadorette, Jules; Lecomte, Roger; Hunting, Darel J.; Pavan, Roberto A.; Zeisler,

Stefan K.; Lier, Johan E.

CORPORATE SOURCE: Sherbrooke Molecular Imaging Centre and Department of

Nuclear Medicine and Radiobiology, Faculty of Medicine and Health Sciences, Universite de Sherbrooke,

Sherbrooke, OC, Can.

SOURCE: European Journal of Nuclear Medicine and Molecular

Imaging (2007), 34(2), 247-258 CODEN: EJNMA6: ISSN: 1619-7070

PUBLISHER: Springer
DOCUMENT TYPE: Journal
LANGUAGE: English

AB In vivo detection of apoptosis is a diagnostic tool with potential clin. applications in cardiol. and oncol. Radiolabeled annexin-V (anxV) is an ideal probe for in vivo apoptosis detection owing to its strong affinity for phosphatidylserine (PS), the mol. flag on the surface of apoptotic cells. Most clin. studies performed to visualize apoptosis have used 99mTc-anxV; however, its poor distribution profile often compromises image quality. In this study, tumor apoptosis after therapy was visualized by positron emission tomog. (PET) using 64Cu-labeled streptavidin (SAv), following pre-targeting of apoptotic cells with biotinvlated anxV. Apoptosis was induced in tumor-bearing mice by photodynamic therapy (PDT) using phthalocyanine dyes as photosensitizers, and red light. After PDT, mice were injected i.v. with biotinylated anxV, followed 2 h later by an avidin chase, and after another 2 h with 64Cu-DOTA-biotin-SAv. PET images were subsequently recorded up to 13 h after PDT. PET images delineated apoptosis in treated tumors as early as 30 min after 64Cu-DOTA-biotin-SAv administration, with tumor-to-background ratios reaching a maximum at 3 h post-injection, i.e., 7 h post-PDT. Omitting the administration of biotinylated anxV or the avidin chase failed to provide a clear PET image, confirming that all three steps are essential for adequate visualization of apoptosis. Furthermore, differences in action mechanisms between photosensitizers that target tumor cells directly or via initial vascular stasis were clearly recognized through differences in tracer uptake patterns detecting early or delayed apoptosis. This study demonstrates the efficacy of a three-step 64Cu pretargeting procedure for PET imaging of apoptosis. These data also confirm the usefulness of small animal PET to evaluate cancer treatment protocols.

CC 8-9 (Radiation Biochemistry)

Copper 64 DOTA biotin streptavidin PET PDT apoptosis; PET imaging annexin V targeted tumor apoptosis photosensitizer

IT Imaging

(tumor; use of pretargeting procedure of phosphatidylserine with biotinylated annexin-V for PET imaging of apoptosis with 64Cu-SAv complex)

IT 956262-96-7P 956428-39-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(use of pretargeting procedure of phosphatidylserine with biotinylated annexin-V for PET imaging of apoptosis with 64Cu-SAv complex)

IT 956262-96-7P

RN

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(use of pretargeting procedure of phosphatidylserine with biotinylated annexin-V for PET imaging of apoptosis with 64Cu-SAv complex)

956262-96-7 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[4-[[5-[3as,4R,6aR]-octahydro-2,5-dioxo-4-cyclopentimidazolyl]]-1-oxopentyl]amino]butyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

~со2Н

REFERENCE COUNT: 45 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 11 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:872573 ZCAPLUS Full-text

DOCUMENT NUMBER: 145:425460 TITLE:

Noninvasive Visualization of in Vivo Drug Delivery of Poly(L-glutamic acid) Using Contrast-Enhanced MRI Ye, Furong; Ke, Tianyi; Jeong, Eun-Kee; Wang, Xuli; AUTHOR(S):

Sun, Yongen; Johnson, Melody; Lu, Zheng-Rong Departments of Pharmaceutics and Pharmaceutical CORPORATE SOURCE:

Chemistry and Radiology, University of Utah, Salt Lake

City, UT, 84108, USA

SOURCE: Molecular Pharmaceutics (2006), 3(5), 507-515

CODEN: MPOHBP; ISSN: 1543-8384

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S):

CASREACT 145:425460

Biomedical imaging is valuable for noninvasive investigation of in vivo drug delivery with polymer conjugates. It can provide real-time information on pharmacokinetics, biodistribution, and drug delivery efficiency of the conjugates. Noninvasive visualization of in vivo drug delivery of polymer conjugates with contrast-enhanced magnetic resonance imaging (MRI) was studied with paramagnetically labeled poly(L-glutamic acid) in an animal tumor model. Poly(L-glutamic acid) is a biocompatible and biodegradable drug carrier for diagnostics and therapeutics. Poly(L-glutamic acid)-1.6-hexanediamine-(Gd-DO3A) conjugates with mol. wts. of 87, 50, and 28 kDa and narrow mol. weight distributions were prepared and studied in mice bearing MDA-MB-231 human breast cancer xenografts. Contrast-enhanced MRI resulted in real-time and three-dimensional visualization of blood circulation, pharmacokinetics, biodistribution, and tumor accumulation of the conjugates, and the size effect on these pharmaceutics properties. The conjugate of 28 kDa rapidly cleared from the circulation and had a relatively lower tumor accumulation. The conjugates with higher mol. wts. exhibited a more prolonged blood circulation and higher tames accumulation. The difference between the conjugates of 87

and 50 kDa was not significant. Contrast-enhanced MRI is effective for noninvasive real-time visualization of in vivo drug delivery of paramagnetically labeled polymer conjugates.

CC 63-5 (Pharmaceuticals)

Section cross-reference(s): 8

IT 912576-20-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(noninvasive visualization of in vivo drug delivery of poly(L-glutamic acid) using contrast-enhanced MRI)

IT 22541-19-1, Gd3+, biological studies 912576-20-60, reaction

products with polyglutamic acid, gadolinium complexes

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(noninvasive visualization of in vivo drug delivery of poly(L-glutamic

acid) using contrast-enhanced MRI)

IT 912576-20-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(noninvasive visualization of in vivo drug delivery of poly(L-glutamic acid) using contrast-enhanced MRI)

RN 912576-20-6 ZCAPLUS

N 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminohexyl)amino]-2-oxoethyl]-, 2,2,2-trifluoroacetate (1:1) (CA INDEX NAME)

CM 1

CRN 889140-15-2 CMF C22 H42 N6 O7

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (noninvasive visualization of in vivo drug delivery of poly(L-glutamic

acid) using contrast-enhanced MRI

REFERENCE COUNT: 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 12 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:836023 ZCAPLUS Full-text

DOCUMENT NUMBER: 147:26007

TITLE: Biodegradable cystamine spacer facilitates the

clearance of Gd(III) chelates in poly(glutamic acid) Gd-DO3A conjugates for contrast-enhanced MR imaging

AUTHOR(S): Ke, Tianyi; Feng, Yi; Guo, Junyu; Parker, Dennis L.;

Lu, Zhena-Rona

Department of Pharmaceutics and Pharmaceutical CORPORATE SOURCE:

Chemistry, University of Utah, Salt Lake City, UT, 84108, USA

SOURCE: Magnetic Resonance Imaging (2006), 24(7), 931-940

CODEN: MRIMDO; ISSN: 0730-725X

PUBLISHER: Elsevier Inc. DOCUMENT TYPE: Journal LANGUAGE: English

> Poly(-glutamic acid) (PGA)-cystamine-[gadolinium (Gd)-DO3A] was prepared in high yield with a high Gd-DO3A conjugation efficiency. Approx. 55% of the carboxylic groups in PGA were loaded with Gd-DO3A via cystamine as the spacer. Cystamine can be readily cleaved by endogenous thiols to release the Gd(III) chelates from the conjugate facilitating Gd(III) excretion after the magnetic resonance imaging (MRI). The contrast-enhanced MRI with PGA-cystamine-(Gd-DO3A) was investigated in mice bearing MDA-MB-231 breast carcinoma xenografts. PGA-1,6-hexanediamine-(Gd-DO3A), a paramagnetic polymer conjugate of a nondegradable spacer, was used as a control. Both conjugates resulted in similar contrast enhancement in the heart, vasculature, liver and kidneys in the first hour post injection. More substantial signal intensity reduction was observed for PGA-cystamine-(Gd- DO3A) in these organs than PGA-1.6hexanediamine-(Gd-DO3A) due to release of the Gd chelates from PGA-cystamine-(Gd-DO3A) after the cleavage of the disulfide spacer by the endogenous thiols. Both conjugates resulted in similar tumor enhancement with approx. 70% increased signal intensity in the tumor periphery and 10-40% increased signal intensity in tumor interstitium. No cross-reaction was observed between PGAcystamine-(Gd-DO3A) and human serum albumin, a plasma protein containing a cysteine residue. PGA-cystamine-(Gd-DO3A) resulted in significantly lower Gd(III) tissue retention than PGA-1,6-hexanediamine-(Gd-DO3A) 10 days after the injection in the mice (P<.05). The conjugation of Gd(III) chelates to biomedical copolymers via the degradable disulfide spacer resulted in significant contrast enhancement in the blood pool and tumor tissue but minimal long-term Gd(III) tissue retention.

CC 8-9 (Radiation Biochemistry)

TТ Imaging

> (tumor; role of biodegradable cystamine spacer in clearance of Gd(III)chelates in polv(glutamic acid)Gd-DO3A conjugates for contrast enhanced magnetic resonance imaging of breast carcinomas)

25513-46-6DP, reaction products with acetic acid tetraazacvclododecane cystamine derivs. 585531-76-6DP, polyglutamic acid derivs., gadolinium complexes 889140-15-2DF, polyglutamic acid derivs., gadolinium complexes

RL: PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(role of biodegradable cystamine spacer in clearance of Gd(III) chelates in poly(glutamic acid)Gd-DO3A conjugates for contrast enhanced magnetic resonance imaging of breast carcinomas)

114873-37-9P 122555-91-3P 485800-28-0P 585531-76-6P TT 889140-15-2P 938041-81-7P

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(role of biodegradable cystamine spacer in clearance of Gd(III)chelates in poly(glutamic acid)Gd-DO3A conjugates for contrast enhanced magnetic resonance imaging of breast carcinomas)

889140-15-2DP, polyglutamic acid derivs., gadolinium complexes

RL: PKT (Pharmacokinetics); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(role of biodegradable cystamine spacer in clearance of Gd(III)chelates in poly(glutamic acid)Gd-DO3A conjugates for contrast enhanced magnetic resonance imaging of breast carcinomas)

889140-15-2 ZCAPLUS RN

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6aminohexyl)amino]-2-oxoethyl]- (CA INDEX NAME)

IΤ 889140-15-2P

> RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation): RACT (Reactant or reagent)

(role of biodegradable cystamine spacer in clearance of Gd(III)chelates in poly(glutamic acid)Gd-DO3A conjugates for contrast enhanced magnetic resonance imaging of breast carcinomas)

RN 889140-15-2 ZCAPLUS

1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-CN aminohexvl)aminol-2-oxoethvll- (CA INDEX NAME)

REFERENCE COUNT:

24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

L80 ANSWER 13 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 2006:779890 ZCAPLUS Full-text 145:391420

Structure-Activity Relationships of 111In- and 99mTc-Labeled Ouinolin-4-one Peptidomimetics as Ligands for the Vitronectin Receptor: Potential Tumor Imaging Agents

Harris, Thomas D.; Kalogeropoulos, Shirley; Nguyen, AUTHOR(S):

Tiffany; Dwyer, Gregory; Edwards, D. Scott; Liu, Shuang; Bartis, Judit; Ellars, Charles; Onthank, Dave;

Yalamanchili, Padmaja; Heminway, Stuart; Robinson, Simon; Lazewatsky, Joel; Barrett, John

CORPORATE SOURCE: Discovery Research, Bristol-Myers Squibb Medical

Imaging, N. Billerica, MA, 01862, USA

SOURCE: Bioconjugate Chemistry (2006), 17(5), 1294-1313

CODEN: BCCHES: ISSN: 1043-1802

PUBLISHER: American Chemical Society Journal

DOCUMENT TYPE:

LANGUAGE: English

AB The integrin receptor $\alpha v\beta 3$ is overexpressed on the endothelial cells of growing tumors and on some tumor cells themselves. Radiolabeled $\alpha v \beta 3$ antagonists have demonstrated potential application as tumor imaging agents and as radiotherapeutic agents. This report describes the total synthesis of eight new HYNIC and DOTA conjugates of receptor av83 antagonists belonging to the quinolin-4-one class of peptidomimetics, and their radiolabeling with 99mTc (for HYNIC) and 111In (for DOTA). Tethering of the radionuclidechelator complexes was achieved at two different sites on the quinolin-4-one mol. All such derivs, maintained high affinity for receptor $\alpha v \beta 3$ and high selectivity vs. receptors aIIbβ3, avβ5, a5β1. Biodistribution of the radiolabeled compds. was evaluated in the c-neu Oncomouse mammary adenocarcinoma model. DOTA conjugate 111In-TA138 presented the best biodistribution profile. Tumor uptake at 2 h postinjection was 9.39% of injected dose/g of tissue (%ID/g). Activity levels in selected organs was as follows: blood, 0.54% ID/g; liver, 1.94% ID/g; kidney, 2.33% ID/g; lung, 2.74% ID/q; bone, 1.56% ID/q. A complete biodistribution anal. of 111In-TA138 and the other radiolabeled compds. of this study are presented and discussed. A scintigraphic imaging study with 111In-TA138 showed a clear delineation of the tumors and rapid clearance of activity from nontarget tissues. 8-9 (Radiation Biochemistry)

CC

prepn radiolabeled quinolinone peptidomimetic vitronectin receptor tumor imaging

Scintigraphic agents

Scintigraphy

Structure-activity relationship

(SAR and preparation of 111In- and 99mTc-labeled guinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

Vitronectin receptors

RL: BSU (Biological study, unclassified); BIOL (Biological study) (SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

Mammary gland, neoplasm

(adenocarcinoma; SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

Carcinoma

(mammary adenocarcinoma; SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tamor imaging agents)

Pharmacokinetics

(organ uptake; SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

Imaging

(tumer; SAR and preparation of 111In- and 99mTc-labeled

quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents) ${}^{\circ}$

IT Integrins

RL: BSU (Biological study, unclassified); BIOL (Biological study)
(aIID\$3, SAR and preparation of 111In- and 99mTc-labeled
quinolin-4-one peptidomimetics as ligands for vitronectin receptor and
potential tumor imaging agents)

IT Integrins

RL: BSU (Biological study, unclassified); BIOL (Biological study) ($\alpha v \beta 3$; SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

IT Integrins

RL: BSU (Biological study, unclassified); BIOL (Biological study) (avp5; SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

IT Integrins

RL: BSU (Biological study, unclassified); BIOL (Biological study) (α5β1; SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

IT 15750-15-9DP, Indium 111, conjugates, biological studies 278172-91-1P
278172-95-5P 278172-98-8P 278172-99-9P 378784-45-3DP, Technetium
99m, conjugates, biological studies 498575-44-3DP, technetium-99 complex
498575-49-8DP, technetium-99 complex 498575-53-4DP, technetium-99
complex 911209-04-6P
RL: DCN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic

RL: DGN (Diagnostic use); PKT (Pharmacokinetics); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential townor imaging agents)

T501-53-1 3406-84-6, Biphenyl-4,4'-disulfonyl chloride 4246-51-9
7790-94-5, Chlorosulfonic acid 66414-73-1 72080-83-2, Benzyl
N-(2-aminoethyl)carbamate 77087-60-6 83948-53-2 98541-64-1
114559-25-0 137076-54-1, DOTA tri(tert-butyl) ester 185563-93-3
206055-18-7 208580-23-8 208580-27-2 277315-96-5 277316-23-1
277316-26-4 277316-29-7 848083-49-8 911141-44-1
RI: RCT (Reactant); RACT (Reactant or reagent)

(SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tomor imaging agents)

TT 40324-66-1P 57932-18-0P 220156-99-0P 250612-31-8P 277315-53-4P 277315-71-6P 277315-77-2P 277315-83-0P 277315-84-1P 277315-85-2P 277315-98-8P 277315-98-9P 277315-98-9P 277315-99-9P 277315-97-6P 277316-03-7P 277316-09-3P 277316-00-4P 277316-01-5P 277316-02-6P 277316-03-7P 277316-40-2P 277316-10-6P 277316-11-7P 277316-24-2P 277316-48-9P 277316-50-4P 277316-41-3P 277316-45-8P 277316-45-9P 277316-41-3P 277316-58-2P 911141-45-2P 911141-45-2P 911141-47-4P 911141-48-5P 911141-49-6P 911141-50-9P 911141-52-1P 911141-53-2P 911141-54-3P 911141-54-3P P1141-54-3P P1141-54-3P

(Reactant or reagent)
(SAR and preparation of 111In- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential

tumor imaging agents)
T 911141-45-2DP, technetium-99 complex 911141-55

911141-45-2DP, technetium-99 complex 911141-55-4P 911141-56-5P RL: SPN (Synthetic preparation); PREP (Preparation)

(SAR and preparation of 1111n- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor imaging agents)

IT 277316-46-8P 911141-54-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(SAR and preparation of 1111n- and 99mTc-labeled quinolin-4-one peptidomimetics as ligands for vitronectin receptor and potential tumor immains agents)

RN 277316-46-8 ZCAPLUS

 $\begin{array}{lll} & 1,4,7,10^{-1} & \text{Rorious} \\ & 1,4,7,10^{-1} & \text{Total acayclodode cane-1,4,7-triacetic acid, } & 10-[2-[[3-[3-[[[(2S)-2-carboxy-2-[([2,4,6-trimethylphenyl)sulfonyl]amino]ethyl]amino]ethyl]amino]ethyll-acoxo-1(4H)-quinolinyl]propyl]amino]-2-oxoethyll-, & a.a., & a.trisc(1,1-dimethylethyl) ester (9CI) & (CA INDEX NAME) & (CA$

Absolute stereochemistry.

RN 911141-54-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[3-[[[(2S)-2-carboxy-2-[([2,4,6-trimethylphenyl]sulfonyl]amino]ethyl]amino]carbonyl]-7-[(1H-imidazol-2-ylamino)methyl]-4-oxo-1(4H)-quinolinyl]propyl]amino]-2oxoethyl]-, bis(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 277315-74-9 CMF C45 H61 N11 O13 S

Absolute stereochemistry.

CM 2

CRN 76-05-1

CMF C2 H F3 O2

REFERENCE COUNT: 50 THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 14 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:681369 ZCAPLUS Full-text

DOCUMENT NUMBER: 145:146029

TITLE: Preparation of peptide-containing compounds for

targeting cells expressing NP-1 receptor

INVENTOR(S): Von Wronski, Mathew A.; Marinelli, Edmund R.; Nunn, Adrian D.; Pillai, Radhakrishna; Ramalingam,

Kondareddiar; Tweedle, Michael F.; Linder, Karen E.;

APPLICATION NO

US 2006-342050

A 20060127

Nanjappan, Palaniappa; Raju, Natarajan

PATENT ASSIGNEE(S): Bracco International B.V., Neth.

SOURCE: U.S. Pat. Appl. Publ., 98 pp., Cont.-in-part of Ser.

No. US 2001-871974, CODEN: USXXCO

KIND DATE

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

	PAIENI NO.					KIND DATE				APPL		DAIL						
	2006153775			A1		2006				2006-342050				20060127				
US	2002147136			A1 20021010			US 2001-871974						20010604					
US	7109167				B2 20060919													
WO	2007090022			A2 200708			0809		WO 2	007-		20070125						
WO	2007	2007090022					2007	1122										
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	
		KP,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	
		MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	
		RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,	TR,	TT,	
		TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW							
	RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	
		IS,	ΙT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	
		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,	GH,	
		GM,	KΕ,	LS,	MW,	ΜZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,	
		KG,	ΚZ,	MD,	RU,	ТJ,	TM,	AP,	EA,	EP,	OA							
PRIORITY	APP	LN.	INFO	. :					US 2000-585364						B2 20000602			
										US 2	001-	8719	74		A2 2	0010	604	

OTHER SOURCE(S): MARPAT 145:146029

Other Source(s):

Marker 107140029

The invention provides compds. for targeting endothelial cells, tumor cells or other cells that express the neuropilin-1 (NP-1) receptor, compns. containing the same and methods for their use. The compds. are of the formula A-L-B (A = a monomer, multimer or polymer of TKPPR or analog which specifically binds to NP-1 or cells expressing NP-1 with avidity equal or greater than TKPPR; L = a lipid or a non-lipid (e.g., polymer) linker; B = a substrate). Addn1, the present invention includes diagnostic, therapeutic and radio-therapeutic compns. useful for visualization, therapy or radiotherapy. For example, DPPE-glutaroyl-Gly-Thr-Lys-Pro-Pro-Arg-OH (DPPE-Glu-GTKPPR) was prepared and formulated into gas-filled microbubble compns. for ultrasonic echog. The bubbles bind to human aortic endothelial cells (HAEC) under flow. The number of bubbles bound may increase with time for several minutes at a given flow rate, up to a flow rate producing 1.53 dynes/cm2, while bubbles without the targeting moiety (DPPE-Glu-GTKPPR) may not bind. However, once bound under a lesser flow rate (e.g., 1.53 dynes/cm2), the shear stress on bubbles

containing DPPE-Glu-GTKPPR may be increased to $6.1 \; dynes/cm2$ without dislodging many of the bound bubbles.

INCL 424009340; 530326000

CC 34-3 (Amino Acids, Peptides, and Proteins)
Section cross-reference(s): 1, 8, 63

ST peptide neuropilin receptor endothelium tumor targeting, antitumor angiogenesis inhibitor peptide deriv prepn; gene therapy radiotherapy peptide deriv; ultrasound imaging endothelium neuropilin peptide

Tumor necrosis factors

RL: BSU (Biological study, unclassified); BIOL (Biological study) (human aortic endothelial cells activated by; preparation of peptide-containing

compds. and compns. for targeting cells expressing neuropilin-1 receptor for diagnosis, imaging, and therapy)

IT 100-46-9, Benzenemethanamine, reactions 1155-64-2 1663-39-4 4530-20-5 5681-36-7 7672-27-7 15401-08-8 29022-11-5 33662-26-9 71989-26-9 71989-35-0 76931-93-6 82911-69-1 106392-12-5 120791-76-6 129223-22-9 166108-71-0 167393-62-6 169543-81-1 198139-51-4 251450-64-3 283176-26-1 377087-81-5D, resin bound 377087-83-7D, resin-bound 47044-40-7 897930-81-3 RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of peptide-containing compds. and compns. for targeting cells expressing neuropilin-1 receptor for diagnosis, imaging, and therapy)

IT 897930-81-3
 RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of peptide-containing compds. and compns. for targeting cells expressing neuropilin-1 receptor for diagnosis, imaging, and therapy)

RN 897930-81-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[3-[[5-(2,5-diydro-2,5-dioxo-1H-pyrrol-1-yl)-1-oxopentyl]amino]propyl]amino]-2-oxoethyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

L80 ANSWER 15 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:542454 ZCAPLUS Full-text

DOCUMENT NUMBER: 145:34213

TITLE: MRI-guided photodynamic therapy for cancer

INVENTOR(S): Lu, Zheng-Rong, Viadya, Anagha; Ke, Tianyi
PATENT ASSIGNEE(S): University of Utah Research Foundation, USA

SOURCE: PCT Int. Appl., 34 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

LANGUAGE: Engl FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	TENT :				KIND DATE				APPL											
WO	2006	0607	97				20060608													
WO	W: AE, AG, AL,								BB	B.C	BB	PW.	BV	B7.	CA	CH				
							DE,													
							ID,													
							LT,													
							NZ,													
							TJ,													
					ZM,		10,	111,	114,	111,	,	10,	OH,	00,	00,	04,	٧٠,			
	PW.						CZ,	DE	DK	EE	ES	FT	FR	CB	CR	нп	TE			
							MC,													
							GN,													
							NA.													
					RU.			,	~=,	~-,	,	00,			,	,	,			
IIA	2005							0608		AII 2	005-		20051202							
	2589																			
										EP 2005-853048										
							CZ,													
							LV,													
KR	2007																			
PRIORIT										US 2004-633255P										
										WO 2										

AB Disclosed is a method of therapy used in combination with a diagnostic tool for enhanced photodynamic therapy using MRI, called (magnetic resonance imaging)-guided photodynamic therapy. The methods of the present invention include administration of MRI contrast agent-labeled polymer photosensitizer conjugates, detection and localization of tumor or cancer tissues with contrast-enhanced MRI and specific illumination and treatment of localized target tissues, such as tumors or cancer cells, using laser energy. The delivered laser energy activates the photosensitizer accumulated in the target tissue, resulting in treatment. Also disclosed are novel conjugate compds., such as PLGA-Mede-DOTA-Gd complexes, having multi-functionality in that the

complex may include an MRI contrasting agent linked to a photosensitizing agent.

CC 63-6 (Pharmaceuticals)

Section cross-reference(s): 8

ST polyglutamate photosensitizer MRI contrast agent delivery tumor;

photodynamic therapy MRI imaging breast cancer

IT Antitumor agents

Human

Neoplasm

Photodynamic therapy

Photosensitizers, pharmaceutical

(delivery systems for MRI-guided photodynamic therapy of cancer)

T 688-74-6DP, reaction products with polyglutamic acids and DOTA, gadolinium complexes 7440-54-2DP, Gadolinium, reaction products with polyglutamic acids, DOTA, and Mce6 25014-27-1DP, deprotected, pyrrolidone esters, DOTA/porphine gadolinium complexes 25038-53-3DP, deprotected, pyrrolidone esters, DOTA/porphine derivs., gadolinium complexes 869140-15-2DP, reaction products with polyglutamic acids,

gadolinium complexes

RL: DGN (Diagnostic use); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(delivery systems for MRI-guided photodynamic therapy of cancer)

IT 889140-15-2DF, reaction products with polyglutamic acids, gadolinium complexes

RL: DGN (Diagnostic use); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(delivery systems for MRI-guided photodynamic therapy of cancer)

RN 889140-15-2 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminchexvl)aminc]-2-oxoethyl]- (CA INDEX NAME)

L80 ANSWER 16 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2006:343390 ZCAPLUS Full-text

DOCUMENT NUMBER: 144:398254

TITLE: Targeted imaging and/or therapy using the Staudinger

ligation

INVENTOR(S): Robillard, Marc S.; Gruell, Holger

PATENT ASSIGNEE(S): Koninklijke Philips Electronics N.V., Neth.

SOURCE: PCT Int. Appl., 59 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

	PATENT NO.						KIND		DATE APPLICATION NO.												
	WO	2006			20060413 20060713				2005-IB53258				20051004								
		W:	AE,	AG.	AL.	AM.	AT.	AU,	AZ.	BA.	BB.	BG.	BR.	BW.	BY.	BZ.	CA,	CH,			
								DE,													
								ID,													
								LU,													
			NA.	NG.	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT.	RO,	RU,	SC,	SD,	SE,	SG,			
			SK,	SL,	SM,	SY,	TJ.	TM.	TN,	TR.	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,			
			YU,	ZA,	ZM,	ZW															
		RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,			
			IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,			
			CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,			
			GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,			
			KG,	KZ,	MD,	RU,	TJ,	TM													
	EΡ	1799	273			A2		2007	0627	0627 EP 2005-788346							20051004				
		R:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,			
			IS,	IT,	LI,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR				
	CN 101068577				A		20071107 CN 2005-80034471							20051004							
	IN 2007CN01400					A		2007	0831		IN 2007-CN1400						20070405				
PRIOR	PRIORITY APPLN. INFO.:										EP 2004-104913					A 2	0041	007			
											WO 2	005-	IB53:	258	1	vi 2	0051	004			

OTHER SOURCE(S):

MARPAT 144:398254

AB The use of a selective chemical and bioorthogonal reaction providing a covalent ligation such as the Staudinger ligation (reaction between an azide and a phosphine), in targeted mol. imaging and therapy is presented, more specifically with interesting applications for pre-targeted imaging or therapy. Current pre-targeted imaging is hampered by the fact that it relies solely on natural/biol. targeting constructs (i.e. biotin/streptavidin). Size considerations and limitations associated with their endogenous nature severely limit the number of applications. The present invention describes how the use of an abiotic, bio-orthogonal reaction which forms a stable adduct under physiol. conditions, by way of a small or undetectable bond, can overcome these limitations. As an example of pre-targeted imaging, injection of a targeting probe comprising a somatostatin receptor-binding peptide linked to an azide is followed by a secondary radiolabeled probe linked to a Staudinger phosphine group. Following in vivo Staudinger ligation, the radiolabel enables detection of the presence of somatostatin receptor-pos. tissue such as neuroendocrine tumor.

CC 63-5 (Pharmaceuticals)

Section cross-reference(s): 1, 8, 21

57260-73-8P 137076-54-1P 149299-82-1P 153086-78-3P 175854-39-4P 192635-89-5P 251564-45-1P 299173-24-3P 361154-31-6P 726698-17-5P 868394-26-7P 882518-79-8P 882518-80-1P 882518-81-2P 882518-82-3P 882518-83-4P 882518-85-6P 882518-86-7P 882518-88-9P

882518-89-0P 882518-90-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(targeted imaging and/or therapy using Staudinger ligation) ΤТ 882518-83-4P 832518-85-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(targeted imaging and/or therapy using Staudinger ligation)

882518-83-4 ZCAPLUS RN

1,4,7,10-Tetraazacvclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[3-CN (diphenylphosphino)-4-(methoxycarbonyl)benzoyl]amino]ethyl]amino]-2oxoethyl |-, tris(1,1-dimethylethyl) ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

RN 882518-85-6 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[3-(diphenylphosphino)-4-(methoxycarbonyl)benzoyl]amino]ethyl]amino]-2-oxoethyl]-, mon(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 882518-84-5

CMF C39 H49 N6 O10 P

PAGE 1-A

PAGE 2-A

CM 2

CRN 76-05-1 CMF C2 H F3 O2

SOURCE:

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S):

L80 ANSWER 17 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 2006:79358 ZCAPLUS Full-text

144:156642

Compositions and methods for treating cancer Mayers, George, L.; Lee, David; Chin, Hsiao Ling Oncologic, Inc., USA PCT Int. Appl., 111 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

> PATENT NO. KIND DATE APPLICATION NO. DATE ----A2 20060126 WO 2005-US26248 WO 2006010165 WO 2006010165 A3 20070208 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM US 2006018908 A1 20060126 US 2004-897530 AU 2005265425 A1 20060126 AU 2005-265425 20050725 CA 2572825 A1 20060126 CA 2005-2572825 20050725 EP 1809332 A2 20070725 EP 2005-802465 20050725 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA, HR, MK, YU

PRIORITY APPLN. INFO.:

US 2004-897530 A 20040723 WO 2005-US26248 W 20050725

The invention features compns. and methods for treating or alleviating a AB symptom of cancer. The compns. and methods of the invention direct supra-LDs of radiation, called Hot-Spots, to virtually all cancer cell types. Cancer is treated by administering a step 1 reagent containing a cell-targeting agent linked to a platform building material; a step 3 reagent containing a targeting moiety and an isotope trapping moiety; and a radiolabeled aqueous soluble set 4 reagent. The cell targeting agent augments cellular uptake of the step 1 reagent. The platform building material detaches from the cell targeting agent upon uptake of the step 1 reagent into the cell and forms an aqueous insol. nano-platform to which the targeting moiety of the step 3 reagent binds. Optionally, a step 2 cell-killing reagent is administered to the subject prior to, after or concurrently with the step 3 reagent to relocate the nano-platform into the tumor extracellular matrix. An example of an agent is an anti-EGF-antibody- dextran-3-indoxyl phosphate-phosphoenol pyruvate conjugate.

CC 63-5 (Pharmaceuticals)

Section cross-reference(s): 1, 8, 15

ΙT Drug delivery systems

> (carriers; radiolabeled tumor-targeted antibody carrier conjugates)

Antibodies and Immunoglobulins Galactosides

Glycosides

Porphyrins

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (conjugates; radiolabeled compr-targeted antibody carrier conjugates)

Glycosides TT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

ΙT

```
(glucuronides, conjugates; radiolabeled tumor-targeted
     antibody carrier conjugates)
Drug delivery systems
     (immunoconjugates; radiolabeled tumor-targeted antibody
     carrier conjugates)
Drug delivery systems
      (immunotoxins; radiolabeled tumor-targeted antibody carrier
     conjugates)
Antitumor agents
Human
Radiopharmaceuticals
      (radiolabeled summar-targeted antibody carrier conjugates)
Albumins, biological studies
Antibodies and Immunoglobulins
Lactams
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
      (radiolabeled tumor-targeted antibody carrier conjugates)
62229-50-9, Eqf
RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
      (antibodies to, conjugates; radiolabeled tumor-targeted
      antibody carrier conjugates)
9024-60-6, Ornithine decarboxylase 9024-77-5, Arginine decarboxylase
RL: BSU (Biological study, unclassified); BIOL (Biological study)
      (inhibitors; radiolabeled tumor-targeted antibody carrier
      conjugates)
9073-60-3
RL: BSU (Biological study, unclassified); BIOL (Biological study)
      (radiolabeled tumor-targeted antibody carrier conjugates)
104-87-0 109-97-7, Pyrrol 119-24-4 122-85-0 616-34-2
                                                                                                                 619-44-3
619-66-9 874-60-2 2646-51-7 3068-32-4 4203-49-0 16522-41-1
21442-01-3 30924-93-7 37293-51-9, Aminodextran 38862-25-8
58626-38-3 60239-18-1, Dota 63379-64-6 76470-66-1, Loracarbef
874201-25-9 874201-26-0 874201-36-2 874201-81-7 874201-87-3
RL: RCT (Reactant); RACT (Reactant or reagent)
     (radiolabeled tumor-targeted antibody carrier conjugates)
61449-63-6P 64244-53-7P 66646-88-6P 78658-49-8P 147804-55-5P
214554-43-5P 214554-44-6P 266341-16-6P 266341-19-9P 623907-52-8P 762241-39-4P 847944-61-0P 847944-62-1P 847944-62-2P 874201-13-5P 874201-19-1P 874201-19-1P 874201-19-1P 874201-19-1P
874201-20-4P 874201-21-5P 874201-22-6P 874201-23-7P 874201-24-8P
874201-27-1P 874201-28-2P 874201-29-3P 874201-30-6P 874201-31-7P
874201-32-8P 874201-33-9P 874201-34-0P 874201-35-1P 874201-37-3P
874201-39-5P 874201-40-8P 874201-41-9P 874201-42-0P 874201-43-1P
874201-44-2P 874201-45-3P 874201-46-4P 874201-47-5P 874201-48-6P
0/4201-49-2F 0/4201-49-3F 0/4201-40-4F 0/4201-49-3F 0/4201-49-3F 0/4201-59-3F 0/4201-59-3F 0/4201-59-3F 0/4201-59-3F 0/4201-59-3F 0/4201-69-4F 0/4201-69-4F 0/4201-69-4F 0/4201-69-4F 0/4201-69-4F 0/4201-69-4F 0/4201-69-4F 0/4201-69-4F 0/4201-79-3F 0/420
874201-77-1P 874201-81-7DP, conjugates with polymer 874201-83-9P
874201-84-0P 874201-85-1P 874201-86-2P 874201-88-4P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
```

59-30-3DP, Folic acid, conjugates 9013-20-1DP, Streptavidin, conjugates 9023-27-2DP, UDP-N-acetylglucosamine enolpyruvyltransferase, conjugates 10098-91-6DP, Yttrium 90, conjugated complexes, biological studies 21442-01-3DP, polymer conjugated derivs. 847944-66-5DP, yttrium 90 complexes 847944-67-6P 847944-68-7P 847944-69-8P 847944-70-1P RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological

(radiolabeled tumor-targeted antibody carrier conjugates)

study); PREP (Preparation); USES (Uses)

(radiolabeled temor-targeted antibody carrier conjugates)

138-08-9D, Phosphoenol pyruvic acid, conjugated derivs. 619-66-9D, 4-Carboxybenzaldehyde, conjugates 9001-78-9D, conjugates 9004-54-0D, Dextran, conjugated derivs. 9031-11-2D, conjugates 13822-19-0D, 3-Indoxyl phosphate, conjugated derivs. 70052-12-9D, q-Difluoromethylornithine, conjugated derivs, 724705-43-5D.

Carbacephem, conjugated derivs. RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (radiolabeled cumor-targeted antibody carrier conjugates)

847944-66-5DP, yttrium 90 complexes RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (radiolabeled tumor-targeted antibody carrier conjugates)

RN 847944-66-5 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[5-[[5-[(3aS, 4S, 6aR) - hexahvdro-2-oxo-1H-thieno[3, 4-d]imidazol-4-v1]-1oxopentyl]amino]pentyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

∼co2H

INVENTOR(S):

L80 ANSWER 18 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 2006:79312 ZCAPLUS Full-text ACCESSION NUMBER:

144:171259 DOCUMENT NUMBER:

TITLE: Preparation of gastrin-releasing peptide compounds for

use in diagnostic imaging or therapy

Cappelletti, Enrico; Lattuada, Luciano; Linder, Karen E.; Marinelli, Edmund; Nanjappan, Palaniappa; Raju, Natarajan; Ramalingam, Kondareddiar; Swenson, Rolf E.;

Tweedle, Michael

PATENT ASSIGNEE(S): Bracco Imaging S.p.A., Italy

SOURCE: U.S. Pat. Appl. Publ., 194 pp., Cont.-in-part of U.S.

Ser. No. 828,925. CODEN: USXXCO

DOCUMENT TYPE: Patent

218

LANGUAGE:

FAMILY ACC. NUM. COUNT: 6

English

PATENT INFORMATION:

	PATENT NO.				KIN									ATE				
US US	2006 2004	0188 1369	30		A1 A1		2006 2004	0126 0715		US 2 US 2	005-	1657	21		2	0050 0030	624	
	7226				B2		2007											
	2004				A2		2004			WO 2	003-	US41	328		2	0031	224	
WO	2004				A3		2004											
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							GD,											
		KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	
		NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SK,	SL,	ТJ,	TM,	TR,	TT,	UA,	
		UG,	US,	UZ,	VN,	YU,	ZW											
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		BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	
		ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	
		TR,	BF,	BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
US	2004	2532	25		A1		2004	1216		US 2	004-	8289	25		2	0040	420	
US	2006	2399	14		A1		2006			US 2	006-	3521	56		2	0060	210	
WO	2007	0025	0.0		A1		2007								2	0060	623	
	W:	AE,	AG,	AL.	AM.	AT.	AU,	AZ.	BA,	BB.	BG.	BR.	BW.	BY.	BZ.	CA.	CH,	
		CN.	co.	CR.	CU.	CZ.	DE,	DK.	DM.	DZ.	EC.	EE.	EG.	ES.	FI.	GB.	GD,	
							HU,											
		KR.	KZ.	LA.	LC.	LK.	LR,	LS.	LT.	LU.	LV.	LY.	MA.	MD.	MG.	MK.	MN.	
							NI,											
		SC.	SD.	SE.	SG.	SK.	SL,	SM.	SY.	TJ.	TM.	TN.	TR.	TT.	TZ.	UA.	UG.	
							ZM.											
	RW:	AT.	BE.	BG.	CH.	CY.	CZ,	DE.	DK.	EE.	ES.	FI.	FR.	GB.	GR.	HU.	IE.	
		IS.	IT.	LT.	LU	LV.	MC,	NL.	PL.	PT.	RO.	SE.	SI.	SK.	TR.	BF.	BJ.	
							GN,											
							NA,											
					RU,													
IN	2006						2007	0706		IN 2	006-	CN23	30		2	0060	626	
	2008						2008			US 2						0070		
PRIORIT	Y APP	LN.								US 2					A2 2			
										WO 2					A2 2			
										US 2					A2 2			
										WO 2						0040		
										US 2					A2 2			
										US 2					A2 2			
OTHER S	OURCE	(S):			MAR	PAT	144:	1712										

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The invention relates to compds. M-N-O-P-G (M is a metal chelator, preferably an Aazta metal chelator or a derivative; N-O-P is a linker containing at least one non- α -amino acid and at least one substituted bile acid; G is the GRP receptor targeting peptide) for use in diagnostic imaging, radiotherapy or phototherapy. Thus, peptide I was prepared and its complex with 177Lu was evaluated for tumor targeting capacity, biodistribution and kinetics in the human PC-3 nude mouse model.

INCL 424001690; 514183000; 534011000

CC 34-3 (Amino Acids, Peptides, and Proteins)

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Section cross-reference(s): 8, 78
55749-98-9P
             55749-99-0P 87096-84-2P, Neuromedin B (swine spinal cord)
422512-72-9P
              422512-75-2P
                             422512-81-0P
                                           721936-47-6P
                                                          721936-49-8P
721936-51-2P
             721936-53-4P
                             721936-55-6P
                                           721936-57-8P
                                                         721936-59-0P
721936-61-4P
             721936-63-6P
                             721936-67-0P
                                           721936-69-2P
                                                         721936-71-6P
721936-73-8P
              721936-75-0P
                             721936-76-1P
                                           721936-78-3P
                                                          721936-80-7P
             721936-92-1P
721936-82-9P
                             721936-94-3P
                                           721936-96-5P
                                                          721936-98-7P
721936-99-8P
             721937-01-5P
                            721937-03-7P
                                           721937-05-9P
                                                          721937-07-1P
721937-09-3P
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              721937-52-6P
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721937-60-6P
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721937-96-8P
              721937-98-0P
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721938-06-3P
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721938-56-3P
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                             722494-02-2P
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808113-04-4P 808113-05-5P 808113-06-6P
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                                                         808113-14-6P
809233-13-4P
             874367-58-5P 874534-72-2P
                                          874534-73-3P
                                                          874537-63-0P
RL: DGN (Diagnostic use); SPN (Synthetic preparation); THU (Therapeutic
use); BIOL (Biological study); PREP (Preparation); USES (Uses)
```

(preparation of gastrin-releasing peptide compds. for use in diagnostic imaging or therapy)

721937-82-2P 721937-90-2P 721937-92-4P 808112-41-6P 808112-74-5P

RL: DGN (Diagnostic use); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of gastrin-releasing peptide compds. for use in diagnostic

imaging or therapy)

RN 721937-82-2 ZCAPLUS CN

L-Methioninamide, N2-[[4-oxo-6-[4-[[4,7,10-tris(carboxymethyl)-1,4,7,10tetraazacvclododec-1-v1|acetv1|-1-piperazinv1|-3(4H)-quinazolinvl|acetv1|-L-qlutaminyl-L-tryptophyl-L-alanyl-L-valylqlycyl-L-histidyl-L-leucyl-

(9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-C

__ SMe

- RN 721937-90-2 ZCAPLUS
- CN L-Methioninamide, N2-[[4-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]-1-piperazinyl]acetyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 2-A

PAGE 2-B

RN 721937-92-4 ZCAPLUS

The thin oin a mide, N2-[[4-[2-[[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetrazacyclododec-l-yl]acetyl]amino]ethyl]-l-piperazinyl]acetyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(CA INDEX NAME)

PAGE 1-B

RN 808112-41-6 ZCAPLUS

CN L-Methioninamide, N2-[4-[[[2-[[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-l-yllacetyllamino]ethyllamino]carbonyllbenzoyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(GA INDEX NAME)

PAGE 1-B

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PAGE 2-A

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PAGE 2-B

- RN 808112-74-5 ZCAPLUS
- CN L-Methioninamide, N2-[4-[bis[2-[[4,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1-yl]acetyl]amino]ethyl]amino]-1,4-dioxobutyl]-L-glutaminyl-L-tryptophyl-L-alanyl-L-valylglycyl-L-histidyl-L-leucyl-(GA INDEX NAME)

PAGE 1-B

$$\sim$$
SMe

HO₂C

PAGE 2-B

L80 ANSWER 19 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2005:1355513 ZCAPLUS Full-text

DOCUMENT NUMBER:

144:99915

TITLE:

INVENTOR(S):

Preparation of lipophilic derivatives of chelate monoamides for use in magnetic resonance imaging Riley, Dennis Patrick; McGhee, William D.

PATENT ASSIGNEE(S): Kereos, Inc., USA SOURCE: PCT Int. Appl., 4

PCT Int. Appl., 47 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT	NO.			KIN	D	DATE			APPL	ICAT	ION :	.00		D	ATE	
					_									-		
WO 200	51228	91		A1		2005	1229		WO 2	005-	US19	966		2	0050	607
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KP,	KR,	KZ,
	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,
	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,
	SL,	SM,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,
	ZA,	ZM,	ZW													
RW	: BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
	AZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,
	RO.	SE.	SI.	SK.	TR.	BF.	BJ.	CF.	CG.	CI.	CM.	GA.	GN.	GO.	GW.	ML.

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MR, NE, SN, TD, TG
    AU 2005253962
                         A1
                               20051229
                                          AU 2005-253962
                                                                  20050607
    CA 2569461
                               20051229
                                           CA 2005-2569461
                                                                  20050607
                         A1
    US 2006008417
                         A1
                               20060112
                                           US 2005-146651
                                                                  20050607
    EP 1768558
                               20070404
                                           EP 2005-757440
                                                                  20050607
                         A1
        R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
            IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR
    JP 2008502726
                         т
                               20080131
                                           JP 2007-527646
                                                                  20050607
PRIORITY APPLN. INFO.:
                                           US 2004-578474P
                                                               P 20040609
                                           US 2004-605180P
                                                               P 20040827
                                           WO 2005-US19966
                                                               W 20050607
OTHER SOURCE(S):
                       CASREACT 144:99915; MARPAT 144:99915
```

AB Compds. useful for associating with nanoparticle or microparticle emulsions to obtain magnetic resonance images permit control of the relaxivity of the signal and readily associate with the particulate components. The compds. are conveniently prepared from achiral derivs. of chelating moieties. Thus, the gadolinium complex of the lipophilic DOTA derivative (I) was prepared in a multistep procedure. This complex was then associated with a nanoparticle/microparticle emulsion and a targeting mol. and used in the magnetic resonance imaging of carcinoma tumors implanted in rabbits.

IC ICM A61B005-055

ICS C07D225-00

for

CC 78-7 (Inorganic Chemicals and Reactions)

Section cross-reference(s): 8

IT 7440-54-2DP, Gadolinium, DOTA monoamide derivative complexes

871560-93-9P 871560-95-1P

RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of lipophilic derivs. of gadolinium-DOTA chelate monoamides

use in magnetic resonance imaging)

T 115265-97-9P 115288-21-6P 201867-18-7P 871560-74-6P 871560-77-9P 871560-80-4P 871560-85-9P

871560-89-3P 871560-91-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of lipophilic derivs. of gadolinium-DOTA chelate monoamides

for use in magnetic resonance imaging)

IT 971560-93-9P 871560-95-1P

RL: DGN (Diagnostic use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

 $(preparation \ of \ lipophilic \ derivs. \ of \ gadolinium-DOTA \ chelate \ monoamides \\ for$

use in magnetic resonance imaging)

RN 871560-93-9 ZCAPLUS

CN Gadolinate(1-), [10-[23-hydroxy-23-oxido-2-(oxo-x0)-11,18,29-trioxo-26-[(1-oxohexadecyl)oxy]-22,24,28-trioxa-3,10,12,19-tetraaza-23-phosphatetratetracont-1-y1]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(4-)-xN1,xN1,xN1,xN10,x01,x0
4,x07]-, hydroqne (9CI) (CA INDEX NAME)

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● H ⁴

PAGE 2-B

- RN 871560-95-1 ZCAPLUS
- Gadolinate(1-), [10-[23-hydroxy-23-oxido-2-(oxo-κ0)-11,18,29-trioxo-CN 26-[(1-oxohexadecv1)oxv]-22.24.28-trioxa-3.10.12.19-tetraaza-23phosphatetratetracont-1-y1]-1,4,7,10-tetraazacyclododecane-1,4,7triacetato(4-)-KN1,KN4,KN7,KN10,KO1,KO 4, KO71- (9CI) (CA INDEX NAME)





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PAGE 1-A

- 871560-77-9P 871560-80-4P 871560-85-9P 871560-89-3P 871560-91-7P
 - RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
- (preparation of lipophilic derivs. of gadolinium-DOTA chelate monoamides for
 - use in magnetic resonance imaging)
- RN 871560-77-9 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[[(1,1dimethylethoxy)carbonyl]amino]hexyl]amino]-2-oxoethyl]-, tris(phenylmethyl) ester (9CI) (CA INDEX NAME)

RN 871560-80-4 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(6-aminohexyl)amino]-2-oxoethyl]-, tris(phenylmethyl) ester (9CI) (CA INDEX NAME)

RN 871560-85-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[[(4-nitrophenoxy)carbonyl]amino]hexyl]amino]-2-oxoethyl]-, tris(phenylmethyl) ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

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- RN 871560-89-3 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[(26R)-23-hydroxy-23-oxido-2,11,18,29-tetraoxo-26-[(1-oxohexadecyl)oxy]-22,24,28-trioxa-3,10,12,19-tetraaza-23-phosphatetratetracont-1-y1]-, tris(phenylmethyl) ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

- RN 871560-91-7 ZCAPLUS
- CN 1,47,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[(26R)-23-hydroxy-23-oxido-2,11,18,29-tetraoxo-26-[(1-oxohexadecyl)oxy]-22,24,28-trioxa-3,10,12,19-tetraaza-23-phosphatetratetracont-1-yl]- (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 20 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2005:1133071 ZCAPLUS Full-text

DOCUMENT NUMBER: 144:57302

AUTHOR(S):

SOURCE:

TITLE: Preparation and Characterization of a

DOTA-Lysine-Biotin Conjugate as an Effector Molecule

for Pretargeted Radionuclide Therapy Hainsworth, James; Harrison, Peter; Mather, Stephen J.

CORPORATE SOURCE: Nuclear Medicine Group, Cancer Research UK, St.

Bartholomew's Hospital, London, UK

Bioconjugate Chemistry (2005), 16(6), 1468-1474

CODEN: BCCHES: ISSN: 1043-1802

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English AB

Pretargeted radionuclide therapy depends on the establishment of a high concentration of secondary binding sites at a tumor to which low-mol. weight radiolabeled effector mols. can be directed. This study describes the simple synthesis of an effector mol. and its subsequent characterization to determine the extent to which it complied with the ideal requirements of such a compound (ε) -DOTA-(α) -biotinamidolysine (DLB) was synthesized in high yield and purity using conventional SPPS methodol. High radiochem, purities were obtained when labeled with several potentially useful radionuclides. The radiolabeled analog bound to streptavidin efficiently with a stoichiometry similar to that of native biotin and showed high stability in serum and upon challenge with acid conditions. Biodistribution studies in normal animals showed a rapid rate of clearance from the blood and low retention of radioactivity by normal

tissues. This design of effector mol. therefore shows promise for further pretargeted radionuclide therapy studies.

63-8 (Pharmaceuticals)

ΙT Antitumor agents

Radiotherapy Stability

(preparation and characterization of a DOTA-lysine-biotin conjugate as an effector mol. for pretargeted radionuclide therapy)

871576-45-3P 871576-46-4F 871576-47-5P 188428-79-7P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and characterization of a DOTA-lysine-biotin conjugate as an effector mol. for pretargeted radionuclide therapy)

871576-46-49

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and characterization of a DOTA-lysine-biotin conjugate as an effector mol. for pretargeted radionuclide therapy)

871576-46-4 ZCAPLUS RN

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-oxo-1H-thieno[3, 4-d]imidazol-4-y1]-1oxopentyl]amino]hexyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B



DOCUMENT NUMBER:

THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 26 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 21 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2005:673150 ZCAPLUS Full-text

143:168816

TITLE: Methods for imaging the lymphatic system using

dendrimer-based contrast agents

INVENTOR(S): Brechbiel, Martin W.; Kobavashi, Hisataka; Chovke, Peter L.; Morris, John C.; Waldmann, Thomas A.

The Government of the United States of America as PATENT ASSIGNEE(S):

Represented by the Secretary of the Department of

Health and Human Services, USA

SOURCE: PCT Int. Appl., 71 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					_	DATE			APPL						ATE	
WO	2005	0679	82							WO 2						0050	
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	2005 1722 R:	RO, MR, 1714 825 AT,	SE, NE, 24	SI, SN,	SK, TD, A1 A2 CH,	TR, TG	BF,	BJ, 0804 1122 DE,	CF,	CG, US 2 EP 2 EE,	CI, 004- 005- ES,	CM, 7569 7224 FI,	GA, 48 40 FR,	GN,	GQ, 2 2 GR,	GW, 0040 0050	ML, 113 112
RIORITY	Y APP	LN.	INFO	. :						US 2						0040	

- AB Methods are disclosed for lymphatic-system imaging using dendrimer conjugates as contrast agents. The disclosed methods are applicable to the imaging of all lymphatic structures, but in particular embodiments are particularly suited for imaging specific parts of the lymphatic system such as lymph nodes or lymphatic vessels. The methods permit the assessment of abnormal conditions within the lymphatic system, such as lymphoma/lymphoproliferative disease, inflammation, and cancer metastasis. The methods also may be used to identify and locate lymph nodes into which lymph fluid flows from a times.
- IC ICM A61K049-00

PR

- CC 8-9 (Radiation Biochemistry)
- 67-43-6D, Dtpa, dendrimer-conjugated complexes 5109-69-3D, Doxa, dendrimer-conjugated complexes 14701-22-5D, Nickel ion (2+), dendrimer-conjugated complexes, biological studies 14913-52-1D. Neodymium ion (3+), dendrimer-conjugated complexes, biological studies 15158-11-9D, Copper ion (2+), dendrimer-conjugated complexes, biological studies 15438-31-0D, Ferrous ion, dendrimer-conjugated complexes, biological studies 16065-83-1D, Chromium ion (3+), dendrimer-conjugated complexes, biological studies 16397-91-4D, Manganese ion (2+), dendrimer-conjugated complexes, biological studies 18472-30-5D, Erbium ion (3+), dendrimer-conjugated complexes, biological studies 18923-27-8D, Ytterbium ion (3+), dendrimer-conjugated complexes, biological studies 20074-52-6D, Ferric ion, dendrimer-conjugated complexes, biological studies 22541-14-6D, Praseodymium ion (3+), dendrimer-conjugated complexes, biological studies 22541-17-9D, Samarium ion (3+), dendrimer-conjugated complexes, biological studies 22541-19-1D, Gadolinium ion (3+), dendrimer-conjugated complexes, biological studies 22541-20-4D, dendrimer-conjugated complexes, biological studies 22541-21-5D, Dysprosium ion (3+), dendrimer-conjugated complexes, biological studies 22541-22-6D, Holmium ion (3+), dendrimer-conjugated complexes, biological studies

22541-53-3D, Cobalt ion (2+), dendrimer-conjugated complexes, biological studies 56491-86-2D, Nota, dendrimer-conjugated complexes 60239-18-1D, Dota, dendrimer-conjugated complexes 60239-22-7D, Teta, dendrimer-conjugated complexes 108414-96-6D, 1b4m, dendrimer-conjugated complexes 113786-33-7D, Bopta, dendrimer-conjugated complexes 114873-3 7-9D, DO 3A, dendrimer-conjugated complexes 120041-08-9D, Hp-do3a, dendrimer-conjugated complexes 149979-17-9D, DO 3MA, dendrimer-conjugated complexes 150467-20-2D, dendrimer-conjugated complexes 160363-61-1D, dendrimer-conjugated complexes RL: DGN (Diagnostic use); BIOL (Biological study); USES (Uses) (imaging the lymphatic system using dendrimer-based contrast agents)

149979-17-9D, DO 3MA, dendrimer-conjugated complexes

150467-20-2D, dendrimer-conjugated complexes RL: DGN (Diagnostic use); BIOL (Biological study); USES (Uses) (imaging the lymphatic system using dendrimer-based contrast agents)

RN 149979-17-9 ZCAPLUS

1,4,7,10-Tetraazacyclododecane-1,4,7-tricarboxylic acid, CN 10-[2-[(2-aminoethvl)amino]-2-oxoethvl]-α, α', α''trimethyl- (9CI) (CA INDEX NAME)

150467-20-2 ZCAPLUS RN

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(2aminoethvl)aminol-2-oxoethvll- (CA INDEX NAME)

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

L80 ANSWER 22 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 2005:238416 ZCAPLUS Full-text 142:303552

> Method and composition for the treatment of cancer by the enzymatic conversion of soluble radioactive toxic precipitates in the cancer

INVENTOR(S): Mavers, George L.; Rose, Samuel; Rose, Lottie

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 78 pp., Cont.-in-part of U.S. Ser. No. 226,288.

CODEN: USXXCO DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2005058652	A1	20050317	US 2004-898585	20040723
US 2003068382	A1	20030410	US 2002-226288	20020822
PRIORITY APPLN. INFO.:			US 2002-226288 A2	20020822
			US 1999-314422 A3	19990518

- AB The invention features compns. and methods for treating or alleviating a symptom of cancer. The compns. and methods of the invention direct supra-LDs of radiation, called Hot-Spots, to virtually all cancer cell types. The compns. comprise a cell-targeting agent (such as an antibody) which augments cellular uptake of the reagent linked to a platform building material by a carrier. The platform building material detaches from the targeting agent upon uptake of the reagent into the cell. Examples of such compns. are: anti-EGFR antibody-dextran-indoxylphosphate- phosphoenolpyruvate conjugate, transferrin-albumin-bis-3-indoxyl glycoside-Loracarbef conjugate, folate-Igporphyrin- difluoromethylornithine conjugate. Above compns. are administered with enzyme conjugates such as β -lactamase-anti-nitroiodophenol antibody, and with radiopharmaceuticals such as 131I-5-iodo-3-indoxvl galactoside.
- IC ICM G01N033-53

ICS G01N033-567; A61K049-00; A61K039-395 INCL 424178100; 530391100; 435007200

- 63-5 (Pharmaceuticals)
- Section cross-reference(s): 1, 8, 15
- ST
- antitemor immunoconjugate immunotoxin radiopharmaceutical enzyme
- ΙT Antitumor agents
- Neoplasm

Nicotinic agonists

Peptidomimetics Radiopharmaceuticals

Radiotherapy

(targeted immunoconjugate radiopharmaceutical compns.)

59-30-3DP, Folic acid, porphyrin-Iq-difluoromethylornithine conjugate 619-66-9DP, 4-Carboxybenzaldehyde, reaction product with ornithine decarboxylase 9001-78-9DP, lactamase conjugate 9013-20-1DP, Streptavidin, UDP-N-Acetylglucosamine enolpyruvyltransferase conjugate 9023-27-2DP, UDP-N-Acetylglucosamine enolpyruvyltransferase, streptavidin 9024-60-6DP, Ornithine decarboxylase, reaction product with carboxybenzaldehyde 9031-11-2DP, lactamase conjugate 9073-60-3DP, galactosidase conjugate 10043-66-0DP, Iodine 131, compds., biological studies 10098-91-6DP, Yttrium 90, conjugated complexes, biological 37293-51-9DP, Aminodextran, antibody conjugate 40704-75-4DP, N-(2-Hydroxypropyl)methacrylamide polymer, crosslinked conjugates 61449-63-6DP, folate-Iq-difluoromethylornithine conjugate 62229-50-9DP, Egf, Loracarbef-polymer conjugates 70052-12-9DP, porphyrin-Ig-folate 76470-66-1DP, Loracarbef, conjugates 847944-58-5DP, antibody-dextran conjugate 847944-59-6DP, antibody-dextran conjugate 847944-60-9DP, antibody-dextran conjugate 847944-61-0DP, albumin-transferrin conjugate 847944-62-1P 847944-64-3DP. EGF-Loracarbef conjugates 847944-66-5DP, vttrium 90 complexes 847944-67-6P 847944-68-7P 847944-69-8P 847944-70-1P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(targeted immunoconjugate radiopharmaceutical compns.)

II 947944-66-5DF, yttrium 90 complexes RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(targeted immunoconjugate radiopharmaceutical compns.)

RN 847944-66-5 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[5-[[5-[3a5,45,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]pentyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

APPLICATION NO. DATE

L80 ANSWER 23 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:14435 ZCAPLUS Full-text

DOCUMENT NUMBER: 142:107822

TITLE: Pharmaceutical composition comprising somatostatin

analog

INVENTOR(S): Lambert, Oliver; Moser, Katrin

KIND DATE

PATENT ASSIGNEE(S): Novartis A.-G., Switz.; Novartis Pharma G.m.b.H.

SOURCE: PCT Int. Appl., 28 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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WO	2005	0008	93		A2		2005	0106		WO 2	004-	EP67	94		2	0040	623
WO	2005	0008	93		A3		2005	0407									
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
		GE.	GH.	GM.	HR.	HU.	ID.	IL.	IN.	IS.	JP.	KE.	KG.	KP.	KR.	KZ.	LC.

(9CI) (CA INDEX NAME)

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                              20050106 CA 2004-2529449
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    EP 1648934
                         A2
                              20060426 EP 2004-740213
                                                                 20040623
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    US 2007093412
                        A1
                              20070426
                                         US 2005-560751
                                                                 20051214
                              20060228 MX 2005-PA13821
    MX 2005PA13821
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    NO 2006000375
                        Α
                              20060124
                                          NO 2006-375
                                                                 20060124
                                                             A 20030624
PRIORITY APPLN. INFO.:
                                          GB 2003-14695
                                          GB 2003-25388
                                                              A 20031030
                                          WO 2004-EP6794
                                                             W 20040623
                        MARPAT 142:107822
OTHER SOURCE(S):
AB
     The present invention describes parenteral pharmaceutical compns. comprising a
     somatostatin analog and novel somatostatin analogs.
    ICM C07K014-655
    ICS A61K038-31; C07K007-06
    2-5 (Mammalian Hormones)
    Section cross-reference(s): 34, 63
TΤ
    Antitumor agents
    Cushing's syndrome
    Drug delivery systems
    Neoplasm
       (pharmaceutical composition comprising somatostatin analog)
    820232-46-0P 820232-47-1P 820232-48-2P
    RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
    (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
    (Uses)
        (pharmaceutical composition comprising somatostatin analog)
    800232-48-2P
    RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
    (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
    (Uses)
       (pharmaceutical composition comprising somatostatin analog)
RN
   820232-48-2 ZCAPLUS
    Cyclo[(2R)-2-phenylqlycyl-D-tryptophyl-L-lysyl-O-(phenylmethyl)-L-tyrosyl-
CN
    L-phenylalanyl-(4R)-4-[[[[2-[[[4,7,10-tris(carboxymethyl)-1,4,7,10-
    tetraazacvclododec-1-vl]acetvl]amino]ethvl]amino]carbonvl]oxv]-L-prolvl]
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PAGE 1-A

PAGE 1-B

L80 ANSWER 24 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 2002:657986 ZCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 137:190759

TITLE: Amino derivatives of biotin and their conjugates with

macrocyclic chelating agents

INVENTOR(S): Paganelli, Giovanni; Chinol, Marco; Ginanneschi, Mauro PATENT ASSIGNEE(S): Sigma-Tau Industrie Farmaceutiche Riunite S.p.A.,

Italy

PCT Int. Appl., 25 pp. SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAI	ENI.	NO.			KIN	D	DATE			APPL	ICAI	TON .	NO.		D.	AIE	
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WO	2002	0660	75		A2		2002	0829		WO 2	002-	IT91			2	0020	215
WO	2002	0660	75		A3		2003	0130									
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            PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG,
            US, UZ, VN, YU, ZA, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
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                        A1 20020829 CA 2002-2436242
    CA 2436242
    AU 2002237517
                        A1
                             20020904 AU 2002-237517
                                                                20020215
    EP 1359943
                        A2
                             20031112 EP 2002-703851
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                        В1
                             20051012
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                       T3 20060316
    ES 2248522
                                         ES 2002-703851
                                                                20020215
                             20040630
    MX 2003PA07317
                       A
                                         MX 2003-PA7317
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    US 2004067199
                       A1 20040408
                                          US 2003-468075
                                                                20030930
PRIORITY APPLN. INFO.:
                                          IT 2001-RM79
                                                            A 20010216
                                          WO 2002-IT91
                                                            W 20020215
                       MARPAT 137:190759
OTHER SOURCE(S):
     Amino biotin derivs, are prepared and used for the preparation of conjugates
AR
     with radionuclides for use in human and animal therapy and diagnostics,
     particularly for the diagnosis and therapy of pathol. conditions such as
     tumors. A reduced biotinylhexamethylenediamine conjugate with DOTA was
     prepared
    ICM A61K051-04
IC
CC
    63-6 (Pharmaceuticals)
    Section cross-reference(s): 8, 26
    Antitumor agents
    Chelating agents
    Diagnostic agents
    Radiopharmaceuticals
    Radiotherapy
       (amino derivs. of biotin and their conjugates with macrocyclic
       chelating agents)
    451478-45-8P
ΙT
    RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
    study); PREP (Preparation); USES (Uses)
        (amino derivs, of biotin and their conjugates with macrocyclic
       chelating agents)
    451478-45-8P
    RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
    study); PREP (Preparation); USES (Uses)
        (amino derivs. of biotin and their conjugates with macrocyclic
       chelating agents)
RN
    451478-45-8 ZCAPLUS
CN
    1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[6-[[5-
     [(3aS, 4S, 6aR)-hexahydro-2-oxo-1H-thieno[3, 4-d]imidazol-4-
    yl]pentyl]amino]hexyl]amino]-2-oxoethyl]- (CA INDEX NAME)
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PAGE 1-A

PAGE 1-B

__CO2H

L80 ANSWER 25 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2002:107368 ZCAPLUS Full-text

DOCUMENT NUMBER: 136:167700

TITLE: Preparation of somatostatin analogues for

pharmaceutical use

INVENTOR(S): Albert, Rainer; Bauer, Wilfried; Bodmer, David; Bruns, Christian; Felner, Ivo; Hellstern, Heribert; Lewis,

Ian; Meisenbach, Mark; Weckbecker, Gisbert; Wietfeld,

Bernhard

PATENT ASSIGNEE(S): Novartis A.-G., Switz.; Novartis-Erfindungen

Verwaltungsgesellschaft m.b.H.; et al.

PCT Int. Appl., 25 pp.

SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAI	ENI	NO.			KIM	D	DAIL			MPPL	ICAI	TON .	NO.		D	HIE	
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WO	2002	0101	92		A2		2002	0207	1	WO 2	001-	EP88	24		2	0010	730
WO	2002	0101	92		A3		2002	0919									
WO	2002	0101	92		A9		2002	1017									
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		GM.	HR.	HU.	TD.	TI	TN.	TS.	JP.	KE.	KG.	KP.	KR.	K7	LC.	LK.	LR.

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                              20070611 TW 2001-90118314 20010726
    TW 282341
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                         A1
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    EP 1307486
                         A2
                              20030507 EP 2001-969555
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        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                              20030701 BR 2001-12859
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                              20040219 JP 2002-515921
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                              20040827 NZ 2001-523836
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    ZA 2003000406
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                                         ZA 2003-406
                                                                 20030115
                       A 20050408 IN 2003-CN143
A 20030319 NO 2003-484
A 20030609 MX 2003-PA991
A1 20050120 US 2003-343288
    IN 2003CN00143
                                                                 20030123
    NO 2003000484
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                                                                20030131
                                          US 2003-343288
    US 2005014686
                                                                 20030826
PRIORITY APPLN. INFO.:
                                          GB 2000-18891
                                                              A 20000801
                                          WO 2001-EP8824
                                                             W 20010730
     The invention provides cyclo[{4-(NH2-C2H4-NH-CO-O-)Pro}-Phq-DTrp-Lys-Tyr(4-
```

AB Benzyl)-Phe] (I) , optionally in protected form, or a pharmaceutically acceptable salt or complex thereof, which has interesting pharmaceutical properties. The ability of I to bind to human somatostatin receptors, inhibit GH release, and decrease IGF-1 plasma levels is exemplified. Pharmaceutical compns. containing the analogs are also claimed.

ICM C07K007-00

CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 2

IT Antitumor agents

(pancreas; preparation of somatostatin analogs for pharmaceutical use)

ΙT Angiogenesis inhibitors

Antidiarrheals

Antitumor agents

Diagnosis Drug delivery systems

(preparation of somatostatin analogs for pharmaceutical use)

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of somatostatin analogs for pharmaceutical use in combination with other drugs)

396091-83-0P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of somatostatin analogs for pharmaceutical use in combination with other drugs)

396091-82-0 ZCAPLUS RN

Cvclo[(2S)-2-phenvlqlvcvl-D-trvptophvl-L-lvsvl-O-(phenvlmethvl)-L-tvrosvl-L-phenylalanyl-(4R)-4-[[[2-[[4,7,10-tris(carboxymethyl)-1,4,7,10tetraazacvclododec-1-vllacetvllamino|ethvllamino|carbonvlloxvl-L-prolvll (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

L80 ANSWER 26 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2001:935597 ZCAPLUS Full-text

DOCUMENT NUMBER: 136:54028

TITLE: Preparation of vitronectin receptor antagonist pharmaceuticals

INVENTOR(S): Rajopadhye, Milind; Barrett, John A.; Carpenter, Alan

P., Jr.; Cheesman, Edward H.; Harris, Thomas D.

PATENT ASSIGNEE(S): Dupont Pharmaceuticals Company, USA

SOURCE: PCT Int. Appl., 449 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: Facent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001098294	A2	20011227	WO 2001-US19794	20010621
WO 2001098294	A3	20030109		

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             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
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             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
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                                            US 2000-213212P
                                                                P 20000621
PRIORITY APPLN. INFO.:
                                            WO 2001-US19794
                                                                W 20010621
OTHER SOURCE(S):
                        MARPAT 136:54028
AB
     Compds. (Q)d-Ln-(Ch)d' (Q is a residue having an indazole-type moiety , d = 1-
     10, d' = 1-100, Ln is a linking group, Ch is a metal-bonding unit) were
     prepared for use in the diagnosis and treatment of cancer. The present
     invention provides novel compds. useful for the treatment of rheumatoid
     arthritis. Thus, 2-[[[4-[4-[[[3-[2-[2-[3-[[6-[[1-aza-2-(2-
     sulfophenyl)vinyl]amino](3-pyridyl)]carbonylamino]propoxy]ethoxy]ethoxy]pr
     opyl]amino]sulfonyl]phenyl]phenyl]sulfonyl]amino]-3-[[1-[3-(indazole-2-
     ylamino)propyl](1H-indazol-5-yl)]carbonylamino]propanoic acid was prepared
     (claimed compound). Syntheses of radiopharmaceticals, e.g.,
     99mTc(VnA) (tricine) (phosphine), where VnA represents the vitronectin receptor
     antagonist, are also described.
     ICM C07D403-00
CC
     34-3 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 8, 28, 63, 78
IT
    Angiogenesis
      Antitumor agents
     Atherosclerosis
     Radiopharmaceuticals
     Rheumatoid arthritis
        (preparation of vitronectin receptor antagonist pharmaceuticals)
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ΙT 5704-04-1DP, Tricine, amino acid derivative, TPPTS technetium-99m complexes 277328-73-1P 277328-74-2P 277328-75-3P 277328-76-4P 277328-78-6P 277328-82-2P 277328-79-7P 277328-80-0P 277328-81-1P 277328-83-3P 277328-84-4P 277328-85-5P 277328-86-6P 277328-87-7P 277328-88-8P 277328-89-9P 277328-90-2P 277328-91-3P 277328-92-4P 277328-93-5P 277328-94-6P 277328-95-7P 277328-96-8P 277328-97-9P 277328-98-0P 277328-99-1P 277329-00-7P 277329-01-8P 277329-02-9P 277329-03-0P 277329-04-1P 277329-05-2P 277329-06-3P 277329-08-5P 277329-09-6DP, technetium-99m, tricine 277329-07-4P tris(m-sulfophenyl)-phosphine complexes 277329-10-9DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277329-11-0DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277329-12-1DP, technetium-99m, tricine tris(m-sulfophenvl)-phosphine 277329-13-2DP, technetium-99m, tricine tris(m-sulfophenyl)phosphine complexes 277329-14-3DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277332-11-3DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 278174-58-6P 278174-59-7P 278174-60-0P 278174-61-1P 278174-62-2P 278174-63-3P 278174-64-4P 278174-65-5P 278174-66-6P 278174-67-7P 278174-68-8P 278174-69-9P 278174-70-2P 278174-71-3P 278177-22-3DP, indium-111-labeled 278177-32-5DP, yttrium-90-labeled RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)

- (preparation of vitronectin receptor antagonist pharmaceuticals)
- RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (preparation of vitronectin receptor antagonist pharmaceuticals)
- RN 277329-03-0 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[4-[4[[[(15)-1-carboxy-2-[[[1-[2-[(3,4,5,6-tetrahydro-2-pyridinyl)amino]ethyl]1H-indazol-5-yl]carbonyl]amino]ethyl]amino]sulfonyl]-3,5-dimethylphenoxy]1-oxobutyl]amino]ethyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

- RN 277329-06-3 ZCAPLUS
- CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[4'-[[(1S)-1-carboxy-2-[[3-((1H-imidazol-2-ylamino)methyl]-1-methyl-1Hindazol-6-yl]carbonyl]amino]ethyl]amino]sulfonyl][1,1'-biphenyl]-4yl]sulfonyl]amino]ethyl]amino]-2-oxoethyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

L80 ANSWER 27 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:420991 ZCAPLUS Full-text

DOCUMENT NUMBER:

133:59098

TITLE:

Preparation of vitronectin receptor antagonist

pharmaceuticals

INVENTOR(S): Rajopadhye, Milind; Harris, Thomas David; Cheesman,

Edward H.

PATENT ASSIGNEE(S): Du Pont Pharmaceuticals Company, USA

SOURCE: PCT Int. Appl., 362 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 8

PATENT INFORMATION:

PAT	TENT :	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
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WO	2000	0354	88		A2		2000	0622		WO 1	999-	US30	312		1	9991:	217
WO	2000	0354	88		A3		2000	1109									
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		NO,	NZ,	PL,	RO,	SG,	SI,	SK,	TR,	UA,	VN,	ZA,	AM,	AZ,	BY,	KG,	KZ,
		MD,	RU,	TJ,	TM												
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		PT,	SE														
US	6322	770			B1		2001	1127		US 1	999-	2812	07		1	9990	330
US	2002	0156	80		A1		2002	0207		US 1	999-	2812	09		1	9990:	330

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    US 6548663
                      B1
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                                                             19990330
    CA 2346935
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                             20000622 CA 1999-2346935
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                             20000703 AU 2000-23715
                                                             19991217
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                                      EP 1999-967442
                                                             19991217
    EP 1140203
                       B1
                            20070523
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                       TR 2001-1775
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    ES 2288040
                       T3 20071216
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    US 2003124120
                      A1 20030703 US 2002-269252
                                                             20021011
    US 2003149262
                      A1 20030807
                                        US 2002-306054
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PRIORITY APPLN. INFO.:
                                        IIS 1998-112829P
                                                         P 19981218
                                                        P 19980331
                                        US 1998-80150P
                                        US 1998-112715P
                                                         P 19981218
                                        IIS 1998-112732P
                                                         P 19981218
                                        US 1998-112831P
                                                         P 19981218
                                        US 1999-281050
                                                         A3 19990330
                                        US 1999-281209
                                                          A3 19990330
                                        WO 1999-US30312
                                                          W 19991217
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OTHER SOURCE(S): MARPAT 133:59098

AB Compds. (Q)d-In-Ch (Q is a residue having an indazole-type moiety , d = 1-10, Ln is a linking group, Ch is a metal-bonding unit) were prepared for use in the diagnosis and treatment of cancer, methods of imaging tumors in a patient, and methods of treating cancer in a patient. The present invention also provides novel compds. useful for monitoring therapeutic angiogenesis treatment and destruction of new angiogenic vasculature. Thus, 2-[[[4-[4-[[3-[2-[2-[3-[[6-[[1-aza-2-(2-sulfopheny])viny]]amino](3-pyridyl))carbonylamino]propoxy]ethoxy]ethoxy]propyl]amino]sulfonyl]phenyl] phenyl]sulfonyl]amino]ro-3-[[1-[3-(indazole-2-ylamino)propyl](IH-indazol-5-yl)]carbonylamino]propancia caid was prepared (claimed compound). Syntheses of radiopharmaceticals, e.g., 99mTc(VnA) (tricine) (phosphine), where VnA represents the vitromectin receptor antagonist are also described.

IC ICM A61K047-48

ICS A61K049-00; A61K051-04

34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 8, 28, 63, 78

IT Angiogenesis

Antitumor agents Atherosclerosis Radiopharmaceuticals Rheumatoid arthritis

(preparation of vitronectin receptor antagonist pharmaceuticals)
IT 5704-04-1DP, Tricine, amino acid derivative, TPPTS technetium-99m complexes

14133-76-7DP, Technetium-99, amino acid derivative, tricine and TPPTS complexes, preparation 63995-70-0DP, TPPTS, amino acid derivative, tricine technetium-99m complexes 277328-73-1P 277328-74-2P 277328-75-3P 277328-76-4P 277328-78-6P 277328-79-7P 277328-80-0P 277328-81-1P 277328-82-2P 277328-83-3P 277328-84-4P 277328-85-5P 277328-86-6P 277328-87-7P 277328-88-8P 277328-89-9P 277328-90-2P 277328-91-3P 277328-92-4P 277328-93-5P 277328-94-6P 277328-95-7P 277328-96-8P 277328-97-9P 277328-98-0P 277328-99-1P 277329-00-7P 277329-01-8P 277329-02-9P 277329-03-9P 277329-04-1P 277329-05-2P 277329-06-3P 277329-07-4P 277329-08-5P 277329-09-6DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277329-10-9DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277329-11-0DP, technetium-99m, tricine tris(m-sulfophenyl)phosphine complexes 277329-12-1DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277329-13-2DP, technetium-99m,

tricine tris(m-sulfophenyl)-phosphine complexes 277329-14-3DP, technetium-99m, tricine tris(m-sulfophenyl)-phosphine complexes 277332-11-3DP, technetium-99m, tricine tris(m-sulfophenvl)-phosphine complexes 278174-58-6P 278174-59-7P 278174-60-0P 278174-61-1P 278174-62-2P 278174-63-3P 278174-64-4P 278174-65-5P 278174-66-6P 278174-67-7P 278174-68-8P 278174-69-9P 278174-70-2P 278174-71-3P 278177-22-3DP, indium-111-labeled 278177-32-5DP, yttrium-90-labeled RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of vitronectin receptor antagonist pharmaceuticals)

T 277329-03-0P 277329-06-3P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of vitronectin receptor antagonist pharmaceuticals)

RN 277329-03-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[4-[4[[[(15)-1-carboxy-2-[[[1-[2-[(3,4,5,6-tetrahydro-2-pyridinyl)amino]ethyl]1H-indazol-5-yl]carbonyl]amino]ethyl]amino]sulfonyl]-3,5-dimethylphenoxy]1-oxobutyl]amino]ethyl]amino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

RN 277329-06-3 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[[2-[[4'[[[(15)-1-carboxy-2-[[3-(1H-imidazol-2-ylamino)methyl]-1-methyl-1Hindazol-6-yl]carbonyl]amino]ethyl]amino]sulfonyl][1,1'-biphenyl]-4yl]sulfonyl]amino]ethyl]amino]-2-oxoethyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

L80 ANSWER 28 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:64531 ZCAPLUS Full-text

DOCUMENT NUMBER:

CORPORATE SOURCE:

SOURCE:

133:39944

TITLE: Synthesis, characterization, and imaging performance

of a new class of macrocyclic hepatobiliary MR

contrast agents

AUTHOR(S): Marinelli, Edmund R.; Neubeck, Richard; Song, Bo;

Wagler, Thomas; Ranganathan, Ramachandran S.;

Sukumaran, Kozikhott; Wedeking, Paul W.; Nunn, Adrian; Runge, Val M.; Tweedle, Michael F.

Bracco Research USA, Princeton, NJ, 08540, USA

Investigative Radiology (2000), 35(1), 8-24

CODEN: INVRAV: ISSN: 0020-9996

PUBLISHER: Lippincott Williams & Wilkins

DOCUMENT TYPE: Journal

LANGUAGE: English

RATIONALE AND OBJECTIVES. To investigate the effect of substituent

lipophilicity, substituent position, and overall charge on the hepatobiliary clearance and tolerance of a series of aromatic ring-containing macrocyclic Gd chelates to select a candidate compound for evaluation as a hepatobiliary imaging agent. METHODS. Hepatobiliary clearance was studied in rats. Tissue distribution and tolerance were studied in mice. Imaging was performed in cats, rabbits, and Rhesus monkeys using T1-weighted pulse sequences or T1weighted breath-hold pulse sequences. RESULTS. All the compds. were excreted bimodally. Gd-2,5-BPA-DO3A was found to have the optimal combination of hepatobiliary clearance (47% in rats, 29% in mice) and tolerance (min. LD 5.0

mmol/kg). Initial imaging studies in cats demonstrated the feasibility of Gd-2/5-BPA-DO3A for hepatic imaging. In rabbits with implanted VX-2 adenocarcinoma as a model for metastatic liver disease, Gd-2,5-BPA-DO3A provided sustained hepatic signal intensity (SI) enhancement and lesion conspicuity over a 120-min imaging time course. In Rhesus monkeys with normal liver function, Gd-2,5-BPA-DO3A afforded sustained hepatic SI enhancement and a time-dependent increase in gallbladder SI over the entire 90-min imaging time course. CONCLUSIONS. Gd-2,5-BPA-DO3A provides dramatic and sustained SI enhancement of hepatic tissue in cats, rabbits, and Rhesus monkeys that was superior in all respects to the extracellular space MRI agent, Gd-HP-DO3A, that was employed as a control.

CC 8-9 (Radiation Biochemistry)

Section cross-reference(s): 78

IT Imaging

(temor; synthesis, characterization, and imaging performance of macrocyclic Gd chelates as hepatobiliary MR contrast agents)

IT 7440-54-2DP, Gadolinium, complexes, biological studies 173526-55-1P
173526-57-3P 173526-61-9P 173526-63-9P 173526-70-0P 173526-77-7P
173526-81-3P 275801-54-2P 275801-55-3P 275801-56-4P
275801-57-5P

RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(synthesis, characterization, and imaging performance of macrocyclic Gd chelates as hepatobiliary MR contrast agents)

ΙT	84827-47-4P	167867-06-3P	173526-69-7P	173526-80-2P	173526-83-5P
	173526-84-6P	173526-85-7P	173526-86-8P	173526-88-0P	173526-89-1P
	173526-90-4P	173526-91-5P	173526-93-7P	173526-94-8P	173527-03-2P
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis, characterization, and imaging performance of macrocyclic Gd chelates as hepatobiliary MR contrast agents)

IT 275801-57-5P

RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(synthesis, characterization, and imaging performance of macrocyclic Gd chelates as hepatobiliary MR contrast agents)

RN 275801-57-5 ZCAPLUS

kappa.N10,κ01,κ04,κ07]-, sodium (9CI) (CA INDEX NAME)

Na+

IT 173527-05-4P 275371-67-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis, characterization, and imaging performance of macrocyclic Gd chelates as hepatobiliary MR contrast agents)

RN 173527-05-4 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[(4-cyclohexylphenyl) [2-[(2-(1,1-dimethylethoxy)-2-oxoethyl] (phenylmethyl) amino]ethyl]amino]-2-oxoethyl]-, tris(1,1-dimethylethyl) ester (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

RN 275371-67-0 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-{2-[(4cyclohexylphenyl) [2-[[2-(1,1-dimethylethoxy)-2oxoethyl] [phenylmethyl) aminojethyl] aminoj-2-oxoethyl]- (CA INDEX NAME)

REFERENCE COUNT: 66 THERE ARE 66 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L80 ANSWER 29 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1999:401701 ZCAPLUS Full-text

DOCUMENT NUMBER: 131:55892

TITLE: DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol INVENTOR(S): Griffiths, Gary L.; Hansen, Hans; Govindan, Serengulam

V.

PATENT ASSIGNEE(S): Immunomedics, Inc., USA

SOURCE: PCT Int. Appl., 31 pp.
CODEN: PIXXD2

CODEN: PIX
DOCUMENT TYPE: Patent
LANGUAGE: English

LANGUAGE: Eng FAMILY ACC. NUM. COUNT: 19

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9930745	A2 199906	24 WO 1998-US26579	19981215
WO 9930745	A3 200001	13	
W: AL, AM, AT,	AU, AZ, BA, B	B, BG, BR, BY, CA, CH, CN	CU, CZ, DE,
DK, EE, ES,	FI, GB, GE, H	U, ID, IL, IN, IS, JP, KE	KG, KP, KR,
KZ, LC, LK,	LR, LS, LT, L	U, LV, MD, MG, MK, MN, MW	MX, NO, NZ,
PL, PT, RO,	RU, SD, SE, S	G, SI, SK, SL, TJ, TM, TR	TT, UA, UG,
US, UZ, VN,	YU, ZW		
RW: GH, GM, KE,	LS, MW, SD, S	Z, UG, ZW, AT, BE, CH, CY	DE, DK, ES,
FI, FR, GB,	GR, IE, IT, L	U, MC, NL, PT, SE, BF, BJ	CF, CG, CI,
CM, GA, GN,	GW, ML, MR, N	E, SN, TD, TG	
US 6120768	A 200009	19 US 1997-990843	19971215

AU 9918258 A 19990705 AU 1999-18258 19981215 PRIORITY APPLN. INFO.: US 1997-990843 A1 19971215 US 1993-62662 B1 19930517 US 1995-409960 A2 19950323 US 1995-486166 B2 19950607 US 1996-688781 A2 19960731 WO 1998-US26579 W 19981215

OTHER SOURCE(S): MARPAT 131:55892

AB A radionuclide-chelator conjugate composition for detecting and/or treating lesions in a patient in a pre-targeting protocol comprises pre-targeting the target cell, tissue, or pathogen with a substrate, using a targeting protein that specifically binds a marker substance on the target cell, tissue, or pathogen and to which the substrate is directly or indirectly bound; parenterally injecting the detection or therapeutic composition of the invention which comprises a chelate conjugate of biotin, a chelator, and a chelatable detection or therapeutic agent, and allowing the composition to accrete at the targeted cell, tissue, or pathogen; wherein the chelate conjugate is purified by chromatog, after chelate formation, or further comprises a blood transit-modifying linker or addend that is covalently bound within the chelate conjugate, or both, and using the detection or therapeutic agent to detect or treat the targeted cell, tissue, or pathogen.

IC ICM A61K051-00

CC 8-9 (Radiation Biochemistry)

Section cross-reference(s): 28, 63, 78

IT Anti-infective agents

Antimicrobial agents

Antiqumor agents

Cardiovascular agents

Diagnosis Drug targeting

Infection

Neoplasm

Paramagnetic materials

Parasiticides

(DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Antitumor agents

(carcinoma; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Apritumor agents

(glioma; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Antitumor agents

(leukemia; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Antitumor agents

(lymphoma; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Antitumor agents

(melanoma; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Antitumor agents

(myeloma; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

II Antitumor agents

(neuroblastoma; DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

IT Antitumor agents

(sarcoma; DOTA-biotin derivative metal complexes for the rapeutic and diagnostic use using a pre-targeting protocol) $\,$ 153-94-6D, D-Tryptophan, linker between biotin and DOTA 319-78-8D, D-Isoleucine, linker between biotin and DOTA 328-38-1D, D-Leucine, linker between biotin and DOTA 556-02-5D, D-Tyrosine, linker between biotin and DOTA 640-68-6D, D-Valine, linker between biotin and DOTA 673-06-3D, D-Phenylalanine, linker between biotin and DOTA 923-27-3D, D-Lysine, linker between biotin and DOTA 10043-49-9D, Gold-198, complexes with biotin-linked-DOTA conjugates, biological studies 10098-91-6D, Yttrium-90, complexes with biotin-linked-DOTA conjugates, biological studies 13967-65-2D, Holmium-166, complexes with biotin-linked-DOTA conjugates, biological studies 13968-53-1D, Ruthenium-103, complexes with biotin-linked-DOTA conjugates, biological studies 13981-51-6D, Mercury-197, complexes with biotin-linked-DOTA conjugates, biological studies 14119-09-6D, Gallium-67, complexes with biotin-linked-DOTA conjugates, biological studies 14119-24-5D. Osmium-191, complexes with biotin-linked-DOTA conjugates, biological studies 14133-76-7D, Technetium-99, complexes with biotin-linked-DOTA conjugates, biological studies 14191-64-1D, Praseodymium-142, complexes with biotin-linked-DOTA conjugates, biological studies 14265-75-9D, Lutetium-177, complexes with biotin-linked-DOTA conjugates, biological studies 14265-85-1D, Actinium-225, complexes with biotin-linked-DOTA conjugates, biological studies 14331-95-4D, Ruthenium-105, complexes with biotin-linked-DOTA conjugates, biological studies 14378-26-8D, Rhenium-188, complexes with biotin-linked-DOTA conjugates, biological studies 14391-11-8D, Gold-199, complexes with biotin-linked-DOTA conjugates, biological studies 14391-19-6D, Terbium-161, complexes with biotin-linked-DOTA conjugates, biological studies 14391-96-9D, Scandium-47, complexes with biotin-linked-DOTA conjugates, biological 14687-25-3D, Lead-203, complexes with biotin-linked-DOTA studies conjugates, biological studies 14885-78-0D, Indium-113, complexes with biotin-linked-DOTA conjugates, biological studies 14913-49-6D, Bismuth-212, complexes with biotin-linked-DOTA conjugates, biological studies 14913-89-4D, complexes with biotin-linked-DOTA conjugates, biological studies 14914-68-2D, Antimony-119, complexes with biotin-linked-DOTA conjugates, biological studies 14967-68-1D, Palladium-103, complexes with biotin-linked-DOTA conjugates, biological studies 14981-64-7D, Palladium-109, complexes with biotin-linked-DOTA conjugates, biological studies 14981-79-4D, Praseodymium-143, complexes with biotin-linked-DOTA conjugates, biological studies 14998-63-1D, Rhenium-186, complexes with biotin-linked-DOTA conjugates, biological 15092-94-1D, Lead-212, complexes with biotin-linked-DOTA conjugates, biological studies 15735-74-7D, Platinum-197, complexes with biotin-linked-DOTA conjugates, biological studies 15750-15-9D, Indium-111, complexes with biotin-linked-DOTA conjugates, biological studies 15756-62-4D, Ruthenium-95, complexes with biotin-linked-DOTA conjugates, biological studies 15757-14-9D, Gallium-68, complexes with biotin-linked-DOTA conjugates, biological studies 15757-86-5D, Copper-67, complexes with biotin-linked-DOTA conjugates, biological studies 15758-35-7D, Ruthenium-97, complexes with biotin-linked-DOTA conjugates, biological studies 15760-04-0D, Silver-111, complexes with biotin-linked-DOTA conjugates, biological studies 15765-78-3D, Rhenium-189, complexes with biotin-linked-DOTA conjugates, biological studies 15766-00-4D, Samarium-153, complexes with biotin-linked-DOTA conjugates, biological studies 60239-18-1D, DOTA, biotin-linker conjugates, metal complexes 60239-18-1D, DOTA, biotin-D-amino acid linked 177959-15-8D, linker between biotin and DOTA 227948-63-2D, linker between biotin and DOTA 227948-64-3D, linker between biotin and DOTA 227948-65-4

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

227948-65-4

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(DOTA-biotin derivative metal complexes for therapeutic and diagnostic use using a pre-targeting protocol)

RN 227948-65-4 ZCAPLUS

1,4,7,10-Tetraazacvclododecane-1,4,7-triacetic acid, 10-[2-[[[[5-CN [(3aS, 4S, 6aR)-hexahydro-2-oxo-1H-thieno[3, 4-d]imidazol-4-yl]-1oxopentyl]methylamino]methyl]methylamino]-2-oxoethyl]- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

-co2H

L80 ANSWER 30 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1997:579696 ZCAPLUS Full-text

DOCUMENT NUMBER: 127:228839

TITLE: Pharmaceutical agents containing perfluoroalkyl-

containing metal complexes and the use thereof in tumor therapy and intervention al radiology

INVENTOR(S): Platzek, Johannes; Niedballa, Ulrich; Raduchel, Bernd; Schlecker, Wolfgang; Weinmann, Hanns-Joachim; Frenzel,

Thomas

PATENT ASSIGNEE(S): Schering A .- G., Germany SOURCE:

PCT Int. Appl., 144 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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WO 9730969
                             19970828 WO 1997-EP684
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            MX, NO, NZ, PL, RO, RU, SD, SG, SI, SK, TJ, TM, TR, TT, UA, UG,
        RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                       A1 19970828 DE 1996-19608278 19960223
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    EP 882010
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                            20010502
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
    JP 2000504736
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    AT 200894
                            20010515 AT 1997-903278
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                            20011030 PT 1997-903278
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                            20010130 US 1997-801983
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    TW 477699
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    NO 323547
    GR 3036306
                       T3 20011031
                                        GR 2001-401156
                                                              20010731
                                         DE 1996-19608278 A 19960223
PRIORITY APPLN. INFO.:
                                         US 1996-12506P P 19960229
WO 1997-EP684 W 19970214
```

OTHER SOURCE(S):

MARPAT 127:228839

AB The invention relates to pharmaceutical agents containing perfiuoro alkylated metal complexes RF-L-A and the use thereof in tumor therapy and interventional radiol., in which formula RF is a perfluorinated, straight-chain or branched C chain with the formula -ChF2nX (X = terminal F, Cl, Br, I or H atom and n = 4-30), L is a binding group, and A is a metal complex or the salts thereof of organic and/or inorg, bases or amino acids or amino acid amides. Thus Gd/Dy/Y/Mn complexes of tetraazacyclododecane having amide pendants with perfluoroalkyl groups or polyaminopolycarboxylic acids with pendants containing perfluoroalkyl groups were prepared.

IC ICM C07C229-06

ICS C07C229-76; C07C237-12; C07C311-00; C07D257-02; A61K033-00; C07F001-00; C07F003-00; C07F005-00; C07F007-00

CC 78-7 (Inorganic Chemicals and Reactions)

Section cross-reference(s): 8, 23, 28, 63

ST lanthanide polyaminopolycarboxylate tetraazacyclododecane perfluoroalkyl pendant prepn; tetraazacyclododecane perfluoroalkyl pendant lanthanide manganese prepn; gadolinium polyaminopolycarboxylate tetraazacyclododecane perfluoroalkyl pendant prepn; dysprosium polyaminopolycarboxylate tetraazacyclododecane perfluoroalkyl pendant prepn; yttrium polyaminopolycarboxylate tetraazacyclododecane perfluoroalkyl pendant prepn; polyaminopolycarboxylate tetraazacyclododecane perfluoroalkyl pendant prepn; tumor therapy perfluoroalkyl pendant aza complex; interventional radiol perfluoroalkyl pendant aza complex; interventional radiol

IT Antitumor agents

(rare earth and manganese perfluoroalkyl-containing tetraazacyclododecane and polyaminopolycarboxylate complexes)

IT 195047-10-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(for preparation of rare earth/manganese complexes for use as pharmaceutical

agents in tumor therapy and interventional radiol.)

F 98-59-9, p-Toluenesulfonyl chloride 100-46-9, Benzylamine, reactions

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106-89-8, reactions 107-15-3, 1,2-Ethanediamine, reactions 108-30-5,
Succinic acid anhydride, reactions 108-55-4, Glutaric acid anhydride
110-85-0, Piperazine, reactions 111-26-2, Hexylamine 111-40-0
112-29-8, n-Decyl bromide 112-60-7, Tetraethylene glycol 123-31-9, 1,4-Benzenediol, reactions 143-33-9, Sodium cyanide 294-90-6,
1,4,7,10-Tetraazacyclododecane 307-35-7, Perfluorooctylsulfonyl fluoride
598-21-0, Bromoacetyl bromide 603-35-0, Triphenylphosphine, reactions
647-42-7, 3,3,4,4,5,5,6,6,7,7,8,8,8-Tridecafluoro-1-octanol 678-39-7,
3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,10-Heptadecafluoro-1-decanol
1138-80-3, Benzyloxycarbonylglycine 1738-76-7, Glycine benzyl ester
p-toluenesulfonate 2016-57-1, Decylamine 2043-47-2,
3,3,4,4,5,5,6,6,6-Nonafluoro-1-hexanol 2043-53-0 2043-57-4
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23911-26-4, Diethylenetriaminepentaacetic acid dianhydride 25711-25-5,
N-Benzyloxycarbonylaziridine 30670-30-5, 1H,1H,2H,2H-Perfluorodecylamine
34143-74-3, 1H, 1H, 2H, 2H-Perfluorodecanethiol 38436-14-5,
1-Bromo-3,3,4,4,5,5,6,6,6-nonafluorohexane 38565-52-5
78277-26-6, Benzyl 6-bromohexanoate 78277-30-2, Benzyl
11-bromoundecanoate 114873-37-9 121326-92-9 130147-42-1,
Pentaerythrite monobenzylether 135984-68-8, 2H,2H-Perfluorodecanal
137679-68-6 146432-43-1 168078-14-6 193530-47-1
RL: RCT (Reactant); RACT (Reactant or reagent)
   (for preparation of rare earth/manganese fluoroalkyl-containing
   polyaminopolycarboxylate/tetraazacyclododecane complexes for use as
   pharmaceutical agents in tumor therapy and interventional
   radiol.)
473-25-6P 2991-50-6P 13406-91-2P 50598-29-3P 51740-38-6P
55427-54-8P 89932-70-7P 94190-73-5P 94190-74-6P 113584-32-0P
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147011-35-6P 186095-24-9P 186095-25-0P 186095-26-1P 193528-82-4P
193528-87-9P 193528-89-1P 193528-92-6P 193528-94-8P
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195047-46-2P 195047-47-3P 195047-48-4P 195047-49-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
   (for preparation of rare earth/manganese fluoroalkyl-containing
   polyaminopolycarboxylate/tetraazacyclododecane complexes for use as
   pharmaceutical agents in tomor therapy and interventional
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IT 193528-81-3P 195047-04-2P
 RL: BUU (Biological use, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP

radiol.)

(Preparation); RACT (Reactant or reagent); USES (Uses)
(preparation and demetalation and use as pharmaceutical agent in
tumor therapy and interventional radiol.)

193528-86-8P 193528-88-0P 193528-90-4P 193528-91-5P 193526-93-7P 193529-09-8P 193529-12-3P 193529-16-7P 193529-24-7P 193529-26-9P 193529-28-1P 193529-30-5P 193529-34-9P 193529-36-1P 193529-41-8P 193529-49-6P 193529-46-3P 193529-52-1P 193529-55-4P 193529-57-6P 193530-48-2P 195046-83-4P 195046-84-5P 195046-86-7P 195046-88-9P 195046-95-8P 195046-98-1P 195046-99-2P 195047-02-0P 195047-06-4P 195047-07-5P 195047-08-6P 195047-09-7P 195047-50-8P

RL: BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and use as pharmaceutical agent in tumor therapy and interventional radiol.)

IT 193528-99-3P 193529-01-0P 193529-03-2P 193529-05-4P 195046-90-3P 195046-93-6P

RL: BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and use as pharmaceutical agent in tumor therapy and interventional radiol..)

IT 193528-92-6P 195047-03-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(for preparation of rare earth/manganese fluoroalkyl-containing polyaminopolycarboxylate/tetraazacyclododecane complexes for use as pharmaceutical agents in tumor therapy and interventional radiol.)

RN 193528-92-6 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-(9-ethyl-11,11,12,12,13,13,14,14,15,15,16,16,17,17,18,18,18-heptadecafluoro-10,10-dioxido-2,7-dioxo-10-thia-3,6,9-triazacctadec-1-v1)- (CA INDEX NAME)

RN 195047-03-1 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-triacetic acid, 10-[2-[4-[(heptadecafluorooctyl)sulfonyl]-1-piperazinyl]-2-oxoethyl]- (9CI) (CA INDEX NAME)

IT 193528-93-7P 195047-02-0P

RL: BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and use as pharmaceutical agent in tumor therapy and interventional radiol.)

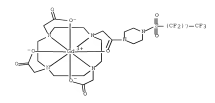
RN 193528-93-7 ZCAPLUS

CN Gadolinium, [10-[9-ethyl-11,11,12,12,13,13,14,14,15,15,16,16,17,17,18,18,1 8-heptadecafluoro-10,10-dioxido-2-(oxo-KO)-7-oxo-10-thia-3,6,9-triazaoctadec-1-y]-1-4,7-10-tetraazacyclododecane-1,4,7-triacetato(3-)-KN1,KN4,KN7,KN10,KO1,KO4,KO7]-

(9CI) (CA INDEX NAME)

RN 195047-02-0 ZCAPLUS

CN Gadolinium, [10-[2-[4-[(heptadecafluorooctyl)sulfonyl]-1-piperazinyl]-2-(0x0-K0)ethyl]-1,4,7,10-tetraazacyclododecane-1,4,7-triacetato(3-)-KN1,KN4,KN7,KN10,K01,K04,K07]-(9CI) (CA INDEX NAME)



L80 ANSWER 31 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1997:184679 ZCAPLUS Full-text

DOCUMENT NUMBER: 126:171905

TITLE:

Somatostatin peptides INVENTOR(S):

Albert, Rainer; Bauer, Wilfried; Bruns, Christian; Chandramouli, Nagarajan; Lewis, Ian; Weckbecker,

Gisbert

PATENT ASSIGNEE(S): Sandoz Ltd., Switz.; Sandoz-Patent-Gmbh;

Sandoz-Erfindungen Verwaltungsgesellschaft M.B.H.; Albert, Rainer; Bauer, Wilfried; Bruns, Christian; Chandramouli, Nagarajan; Lewis, Ian; Weckbecker,

Gisbert

SOURCE: PCT Int. Appl., 32 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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AT 2	10152	T	20011215	AT	1996-924811		19960628
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JP 2	003104998	A	20030409	JP	2002-208012		19960628
SK 2	84087	B6	20040908	SK	1997-1770		19960628
IL 1	22243	A	20050925	IL	1996-122243		19960628
CZ 2	97381	B6	20061115	CZ	1997-4196		19960628
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NO 9	706064	A	19980216	NO	1997-6064		19971223
NO 3	17867	B1	20041227				
US 6	225284	B1	20010501	US	1997-981426		19971229
HK 1	014964	A1	20050408	HK	1999-100124		19990111
PRIORITY .	APPLN. INFO.:			GB	1995-13224	Α	19950629
				GB	1996-429	Α	19960110
				JP	1996-536834	A3	19960628
				WO	1996-EP2840	W	19960628

OTHER SOURCE(S): MARPAT 126:171905

- AB Somatostatin analogs comprising the amino acid sequence -(D/L)Trp-Lys-X1-X2-[X1 = NHCH(CHMCCH2R1)CO [R1 = optionally substituted phenyl) or NHCH(CHR2R2)CO [R2 = ZCH2R1 (X = 0, S), CH2CO2CH2R1, C6H40CH2R1-p, C6H3(CH2R1)OH-3, 4); X2 is an α -amino acid having an aromatic residue on the $C\alpha$ side chain or an amino acid unit selected from Dab, Dpr, Dpm, His, (Bzl)HyPro, thienyl-Ala, cyclohexyl-Ala, and tert-Bu-Ala] or their pharmaceutically acceptable salts or complexes with a detectable element were prepared The Lys residue Lys of the sequence corresponds to the Lys9 residue of native somatostatin-14. Thus, cyclo|HyPro-Phe-DTpr-Lys-Tyr(Bzl)-Phel (I) was prepared by the solid phase method, starting from Fmoc-Phe-SASRIN Resin. IC50 data for binding of I to somatostatin receptor subtypes are tabulated.
- IC ICM C07K014-655 ICS A61K038-31
- CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 1
- ST somatostatin peptide prepn pharmacol property; receptor binding somatostatin peptide; gastric acid secretion somatostatin peptide; antitumor somatostatin peptide; angiogenesis somatostatin peptide; allograft somatostatin peptide; angioplasty somatostatin peptide
- IT Angiogenesis
 - Antitumor agents
- (preparation and pharmacol. properties of somatostatin peptides)
 II 50-99-7, D-Glucose, reactions 141-46-8, Hydroxyacetaldehyde
- 15186-48-8, 2,3-0-Isopropylidene-D-glyceraldehyde 35661-40-6D, resin-bound 57260-73-8 69645-57-4 122350-59-8 134751-65-8 150629-67-7 187223-07-0
 - RL: RCT (Reactant); RACT (Reactant or reagent)
- (preparation and pharmacol. properties of somatostatin peptides)
- RL: RCT (Reactant); RACT (Reactant or reagent)
 - (preparation and pharmacol. properties of somatostatin peptides)
- RN 187223-07-0 ZCAPLUS
- CN Cyclo[L-lysyl-O-(phenylmethyl)-L-seryl-L-phenylalanyl-(4R)-4-[[[2-[[14,7,10-tris(carboxymethyl)-1,4,7,10-tetraazacyclododec-1yl]acetyl]amino]ethyl]amino]carbonyl]oxy]-L-prolyl-L-phenylalanyl-D
 - tryptophyl] (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-C

CO2H

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

L80 ANSWER 32 OF 32 ZCAPLUS COPYRIGHT 2008 ACS on STN 1996:254285 ZCAPLUS Full-text 124:311363

Hydrophilic polymer and radioactive metal complexes as

INVENTOR(S):

locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases Seki, Ikuva; Sato, Toku; Seri, Shigemi; Washino,

Hiroaki

Nihon Mediphysics Co Ltd. Japan PATENT ASSIGNEE(S): SOURCE: Jpn. Kokai Tokkyo Koho, 19 pp.

CODEN: JKXXAF DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
JP 08012597	A	19960116	JP 1993-290080	19931026		
JP 3727074	B2	20051214				
PRIORITY APPLN. INFO.:			JP 1993-290080	19931026		
CT						



AB Biodegradable hydrophilic polymers (polysaccharides and their derivs. containing 1-4 hydrophilic monomer I, with average mol. weight 1 x 103-1 x 106; R1, R2 = H, amino, or hydroxy group; R3 = H, glycol, or carboxymethyl group; R4, R5 = H or hydroxy group) and complex with 1 or >1 radioactive metals are claimed as locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases. Thus, I were prepared and their pharmacokinetics and antitumer and antiinflammatory effects were studied in mice and rats and discussed with their clin. effectiveness.

IC. ICM A61K051-00

CC 8-9 (Radiation Biochemistry)

Section cross-reference(s): 29

hydrophilic polymer radioactive metal complex antitumor;

antiinflammatory hydrophilic polysaccharide radioactive metal complex

ΙT 175783-37-6P 175783-38-7P 175892-38-3DP, complex with indium-111 175892-39-4P 175892-40-7P 176199-54-5P

RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(hydrophilic polymer and radioactive metal complexes as locally administered radio-therapeutic agents for treatment of cancer and

inflammatory diseases) 67-43-6, Diethylenetriamine penta-acetic acid 1398-61-4, Chitin

9012-76-4, Chitosan 10361-82-7, Samarium chloride (SmCl3) 10361-92-9, Yttrium chloride (YCl3) 39271-65-3, Yttrium chloride (90YCl3) 39280-86-9, Glycol chitosan 58259-86-2 149979-17-9, DO 3MA RL: RCT (Reactant); RACT (Reactant or reagent)

(hydrophilic polymer and radioactive metal complexes as locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases)

175783-40-1P 175783-41-2P 175892-38-3P 175892-42-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(hydrophilic polymer and radioactive metal complexes as locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases)

IT 175892-38-30P, complex with indium-111

RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(hydrophilic polymer and radioactive metal complexes as locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases)

RN 175892-38-3 ZCAPLUS

CN Chitosan, 2-hydroxyethyl ether, polymer with 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]-a,a',a''-trimethyl-1,4,7,10-

tetraazacyclododecane-1,4,7-tricarboxylic acid (9CI) (CA INDEX NAME)

CM

CRN 149979-17-9 CMF C21 H40 N6 O7

CM 2

CRN 39280-86-9

CMF C2 H6 O2 . x Unspecified

CM 3

CRN 9012-76-4

CMF Unspecified

CCI PMS, MAN

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

CM 4

CRN 107-21-1

CMF C2 H6 O2

но-си2-си2-он

IT 149979-17-9, DO 3MA

RL: RCT (Reactant); RACT (Reactant or reagent) (hydrophilic polymer and radioactive metal complexes as locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases)

RN 149979-17-9 ZCAPLUS

CN 1,4,7,10-Tetraazacyclododecane-1,4,7-tricarboxylic acid,
10-[2-[(2-aminoethyl)amino]-2-oxoethyl]-a,a',a''trimethyl- [9C1) (CA INDEX NAME)

IT 175892-38-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(hydrophilic polymer and radioactive metal complexes as locally administered radio-therapeutic agents for treatment of cancer and inflammatory diseases)

RN 175892-38-3 ZCAPLUS

CN Chitosan, 2-hydroxyethyl ether, polymer with 10-[2-[(2-aminoethyl)amino]-2-oxoethyl]-a,a',a''-trimethyl-1,4,7,10-tetraazacyclododecane-1,4,7-tricarboxylic acid (9CI) (CA INDEX NAME)

CM

CRN 149979-17-9 CMF C21 H40 N6 O7

CM 2

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CRN 39280-86-9
CMF C2 H6 O2 . x Unspecified

CM 3
CRN 9012-76-4
CMF Unspecified
CCI PMS, MAN

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

CM 4
CRN 107-21-1
CMF C2 H6 O2

HO-CH2-CH2-OH
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L3

=> d his full

(FILE 'HOME' ENTERED AT 08:46:48 ON 21 FEB 2008)

FILE 'ZCAPLUS' ENTERED AT 08:47:31 ON 21 FEB 2008

E US2006-573938/APPS

1 SEA ABB=ON PLU=ON US2006-573938/AP

D SCA

SEL RN

FILE 'REGISTRY' ENTERED AT 08:53:08 ON 21 FEB 2008

65 SEA ABB=ON PLU=ON (118726-52-6/BI OR 17137-11-0/BI OR 294-90-6/BI OR 507475-91-4/BI OR 5292-43-3/BI OR 7429-91-6/BI OR 7439-91-0/BI OR 7439-94-3/BI OR 7440-00-8/BI OR 7440-10-0/BI OR 7440-12-2/BI OR 7440-19-9/BI OR 7440-20-2/BI OR 7440-27-9/B I OR 7440-30-4/BI OR 7440-45-1/BI OR 7440-52-0/BI OR 7440-53-1/ BI OR 7440-54-2/BI OR 7440-60-0/BI OR 7440-64-4/BI OR 7440-65-5 /BI OR 849610-60-2/BI OR 849610-61-3/BI OR 849610-62-4/BI OR 849610-63-5/BI OR 849610-64-6/BI OR 849610-65-7/BI OR 849610-66 -8/BI OR 849610-67-9/BI OR 849610-68-0/BI OR 849610-69-1/BI OR 849610-70-4/BI OR 849610-71-5/BI OR 849610-72-6/BI OR 849610-73 -7/BI OR 849610-74-8/BI OR 849610-75-9/BI OR 849610-76-0/BI OR 849610-77-1/BI OR 849610-78-2/BI OR 849610-79-3/BI OR 849610-80 -6/BI OR 849610-81-7/BI OR 849610-82-8/BI OR 849610-83-9/BI OR 849610-84-0/BI OR 849610-85-1/BI OR 849610-86-2/BI OR 849610-87 -3/BI OR 849610-88-4/BI OR 849610-89-5/BI OR 849610-90-8/BI OR 849610-91-9/BI OR 849610-92-0/BI OR 849610-93-1/BI OR 849610-94 -2/BI OR 849610-95-3/BI OR 849610-96-4/BI OR 849610-97-5/BI OR 849610-98-6/BI OR 849610-99-7/BI OR 849611-00-3/BI OR 849680-88 -2/BI OR 95196-95-5/BI) D SCA

1 SEA ABB=ON PLU=ON L2 AND NRRS>3 D SCA

FILE 'ZCAPLUS' ENTERED AT 09:01:09 ON 21 FEB 2008 L4 1 SEA ABB=ON PLU=ON L3

FILE 'STNGUIDE' ENTERED AT 09:01:27 ON 21 FEB 2008

FILE 'ZCAPLUS' ENTERED AT 09:03:31 ON 21 FEB 2008 L5 11144 SEA ABB=ON PLU=ON L2 (L) PREP/RL L6 1 SEA ABB=ON PLU=ON L5 AND L1 D SCA SEL HIT RN

FILE 'REGISTRY' ENTERED AT 09:04:56 ON 21 FEB 2008

46 SEA ABB—ON PLU—ON (118726-52-6/BI OR 17137-11-0/BI OR 507475-91-4/BI OR 849610-60-2/BI OR 8496610-61-3/BI OR 849610-64-4/BI OR 849610-65-7/BI OR 849610-64-6/BI OR 849610-65-7/BI OR 849610-66-6/BI OR 849610-65-7/BI OR 849610-66-0/BI OR 849610-66-0/BI OR 849610-67-9/BI OR 849610-69-0/BI OR 849610-79-6/BI OR 849610-79-6/BI OR 849610-79-3/BI OR 849610-79-3/BI OR 849610-73-7/BI OR 849610-74-8/BI OR 849610-79-9/BI OR 849610-79-3/BI OR 849610-78-9/BI OR 849610-79-3/BI OR 849610-80-6/BI OR 849610-74-8/BI OR 849610-82-8/BI OR 849610-80-6/BI OR 849610-81-7/BI OR 849610-82-8/BI OR 849610-80-9/BI OR 849610-80-8/BI OR 849610-80-9/BI OR 849610-80-9/BI OR 849610-80-8/BI OR 849610-80-8/BI OR 849610-80-8/BI OR 849610-80-8/BI OR 849610-80-8/BI OR 849610-91-9/BI OR 849610-92-0/BI OR 849610-91-9/BI OR 849610-92-0/BI OR 849610-93-7/BI OR 849610-90-8/BI OR 849610-91-9/BI OR 849610-92-0/BI OR 849610-93-7/BI OR 849610-90-3/BI OR 849610-91-3/BI OR 849610-93-3/BI OR 849610-91-3/BI OR 849610-93-3/BI OR 849610-95-3/BI OR 849610-93-3/BI OR 849610-95-3/BI OR 849610-93-3/BI OR 849610-95-3/BI OR 849610-9

-5/BI OR 849610-98-6/BI OR 849610-99-7/BI OR 849611-00-3/BI OR 849680-88-2/BI OR 95196-95-5/BI)

```
FILE 'ZCAPLUS' ENTERED AT 09:05:08 ON 21 FEB 2008
1.8
            76 SEA ABB=ON PLU=ON L7
1.9
               ANALYZE PLU=ON L8 1- RN HIT : 46 TERMS
    FILE 'REGISTRY' ENTERED AT 09:05:33 ON 21 FEB 2008
             1 SEA ABB=ON PLU=ON 17137-11-0
L10
               D SCA
            45 SEA ABB=ON PLU=ON L7 NOT L10
    FILE 'ZCAPLUS' ENTERED AT 09:05:59 ON 21 FEB 2008
L12
             6 SEA ABB=ON PLU=ON L11
               D SCA
    FILE 'REGISTRY' ENTERED AT 09:06:41 ON 21 FEB 2008
1.13
             0 SEA ABB=ON PLU=ON 507475-91-4P
             1 SEA ABB=ON PLU=ON 507475-91-4
L14
               D SCA
             0 SEA ABB=ON PLU=ON 95196-95-5P
L15
             1 SEA ABB=ON PLU=ON 95196-95-5
L16
               D SCA
            43 SEA ABB=ON PLU=ON L11 NOT (L14 OR L15 OR L16)
    FILE 'ZCAPLUS' ENTERED AT 09:07:26 ON 21 FEB 2008
1.18
             2 SEA ABB=ON PLU=ON L17
               D SCA
    FILE 'REGISTRY' ENTERED AT 09:16:14 ON 21 FEB 2008
T.19
               STRUCTURE UPLOADED
               D SCA L17
T.20
               STRUCTURE UPLOADED
L21
               STRUCTURE UPLOADED
L22
            50 SEA SSS SAM L21
              D SCA
               STRUCTURE UPLOADED
L23
L24
           17 SEA SSS SAM L23
L25
              STRUCTURE UPLOADED
L26
           50 SEA SSS SAM L25
L27
              STRUCTURE UPLOADED
L28
             4 SEA SSS SAM L27
               D SCA
               D STAT OUE L28
               D STAT QUE L26
               D STAT OUE L26
1.29
          2020 SEA SSS FUL L25
               SAVE TEMP L29 PAG938STR25L/A
1.30
             4 SEA SUB=L29 SSS SAM L27
L31
            62 SEA SUB=L29 SSS FUL L27
               SAVE TEMP L31 PAG938STR27L/A
    FILE 'ZCAPLUS' ENTERED AT 09:46:30 ON 21 FEB 2008
             9 SEA ABB=ON PLU=ON L31
   FILE 'REGISTRY' ENTERED AT 09:47:04 ON 21 FEB 2008
          47 SEA ABB=ON PLU=ON L31 NOT L17
L33
L34
              STRUCTURE UPLOADED
```

0 SEA SUB=L29 SSS SAM L34

L35

```
10/573938
1.36
           12 SEA SUB=L29 SSS FUL L34
               D SCA
    FILE 'ZCAPLUS' ENTERED AT 09:53:57 ON 21 FEB 2008
             1 SEA ABB=ON PLU=ON L36
L38
             9 SEA ABB=ON PLU=ON L37 OR L32
             1 SEA ABB=ON PLU=ON L38 AND L1
L39
               SEL RN L38
    FILE 'REGISTRY' ENTERED AT 09:54:59 ON 21 FEB 2008
L40
           273 SEA ABB=ON PLU=ON (934183-16-1/BI OR 111119-28-9/BI OR
                137076-54-1/BI OR 14265-75-9/BI OR 15750-15-9/BI OR 15757-14-9/
               BI OR 317809-26-0/BI OR 33507-63-0/BI OR 705283-66-5/BI OR
               901439-51-8/BI OR 901439-89-2/BI OR 901442-07-7/BI OR 901443-47
               -8/BI OR 91037-65-9/BI OR 934183-14-9/BI OR 934183-15-0/BI OR
               934350-78-4/BI OR 934350-82-0/BI OR 934350-86-4/BI OR 934350-87
               -5/BI OR 10098-91-6/BI OR 110880-55-2/BI OR 110880-57-4/BI OR
               111844-19-0/BI OR 112188-16-6/BI OR 115608-61-2/BI OR 118726-52
                -6/BI OR 128009-23-4/BI OR 135702-31-7/BI OR 137184-55-5/BI OR
                137813-35-5/BI OR 13967-64-1/BI OR 13967-65-2/BI OR 13981-25-4/
               BI OR 13981-56-1/BI OR 14119-08-5/BI OR 14119-09-6/BI OR
               14133-76-7/BI OR 141743-95-5/BI OR 14191-64-1/BI OR 14265-85-1/
               BI OR 14687-25-3/BI OR 14809-53-1/BI OR 14834-85-6/BI OR
               14885-78-0/BI OR 148893-10-1/BI OR 14913-49-6/BI OR 14981-79-4/
               BI OR 15065-93-7/BI OR 15757-86-5/BI OR 15765-31-8/BI OR
               15776-20-2/BI OR 161552-03-0/BI OR 17137-11-0/BI OR 174267-75-5
               /BI OR 188982-12-9/BI OR 22541-18-0/BI OR 22541-19-1/BI OR
               267410-13-9/BI OR 29022-11-5/BI OR 294-90-6/BI OR 36849-05-5/BI
                OR 41444-88-6/BI OR 415706-07-9/BI OR 507475-91-4/BI OR
                5292-43-3/BI OR 585531-74-4/BI OR 6066-82-6/BI OR 623575-85-9/B
               I OR 676544-84-6/BI OR 676544-85-7/BI OR 676553-18-7/BI OR
               676553-19-8/BI OR 7087-68-5/BI OR 713520-27-5/BI OR 728914-72-5
               /BI OR 728914-74-7/BI OR 7429-91-6/BI OR 7439-91-0/BI OR
                7439-94-3/BI OR 7440-00-8/BI OR 7440-10-0/BI OR 7440-12-2/BI
               OR 7440-19-9/BI OR 7440-20-2/BI OR 7440-27-9/BI OR 7440-30-4/BI
                OR 7440-45-1/BI OR 7440-52-0/BI OR 7440-53-1/BI OR 7440-54-2/B
               I OR 7440-60-0/BI OR 7440-64-4/BI OR 7440-65-5/BI OR 766529-14-
               0/BI OR 766529-15-1/BI OR 766529-16-2/BI OR 766529-18-4/BI OR
               766529-19-5/BI OR 766529-20-8/BI OR 766529-22-0/BI OR 766529-24
               -2/BI OR 766529-25-3/BI OR 76652
L41
           65 SEA ABB=ON PLU=ON L40 AND L2
L42
            75 SEA ABB=ON PLU=ON L40 AND M/ELS
L43
            57 SEA ABB=ON PLU=ON L42 NOT L41
1.44
            38 SEA ABB=ON PLU=ON L43 NOT (L31 OR L36)
               D SCA
    FILE 'ZCAPLUS' ENTERED AT 09:59:32 ON 21 FEB 2008
1.45
             8 SEA ABB=ON PLU=ON (L41 OR L42) AND L38
    FILE 'REGISTRY' ENTERED AT 10:00:17 ON 21 FEB 2008
L46
           105 SEA ABB=ON PLU=ON L29 AND Y/ELS
L47
               STRUCTURE UPLOADED
L48
             9 SEA SUB=L29 SSS SAM L47
```

203 SEA ABB=ON PLU-ON L49 NOT L50

FILE 'REGISTRY' ENTERED AT 10:06:58 ON 21 FEB 2008

FILE 'ZCAPLUS' ENTERED AT 10:07:02 ON 21 FEB 2008

142 SEA ABB=ON PLU=ON L49 AND M/ELS

345 SEA SUB=L29 SSS FUL L47

L49 L50

L51

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10/573938
L52
            86 SEA ABB=ON PLU=ON L51
L53
                ANALYZE PLU=ON L52 1- RN HIT : 196 TERMS
    FILE 'REGISTRY' ENTERED AT 10:07:38 ON 21 FEB 2008
1.54
       9 SEA ABB=ON PLU=ON L50 AND Y/ELS
                 D SCA
    FILE 'ZCAPLUS' ENTERED AT 10:08:22 ON 21 FEB 2008
L55
             10 SEA ABB=ON PLU=ON L54
    FILE 'REGISTRY' ENTERED AT 10:08:44 ON 21 FEB 2008
            112 SEA ABB=ON PLU=ON L50 AND LNTH/PG
L56
                 D SCA L2
     FILE 'ZCAPLUS' ENTERED AT 10:11:54 ON 21 FEB 2008
L57
             36 SEA ABB=ON PLU=ON L56
             18 SEA ABB=ON PLU=ON L32 OR L37 OR L45 OR L55
L58
       18 38A ABB—ON PLU—ON L32 OR L37 OR L43 OR L57 OR L57
64196 SEA ABB—ON PLU—ON TUMOUR?/BI OR ?TUMOR?/BI
39 SEA ABB—ON PLU—ON L52 AND L60
39 SEA ABB—ON PLU—ON L52 AND L60
L59
L60
L61
         25232 SEA ABB=ON PLU=ON ?SCAFFOLD?/BI
L62
              2 SEA ABB=ON PLU=ON L49 AND L62
L63
                D SCA
L64
              2 SEA ABB=ON PLU=ON (L51 OR L56) AND L62
L65
            40 SEA ABB=ON PLU=ON (L51 OR L56) AND L60
50 SEA ABB=ON PLU=ON L58 OR L64 OR L65
L66
L67
             8 SEA ABB=ON PLU=ON (L64 OR L65) AND L58
            96 SEA ABB=ON PLU=ON GARLICH J?/AU
L68
L69
            49 SEA ABB=ON PLU=ON SUHR R?/AU
L70
           710 SEA ABB=ON PLU=ON PATTERSON M?/AU
L71
             5 SEA ABB=ON PLU=ON L68 AND (L69 OR L70)
             4 SEA ABB=ON PLU=ON L69 AND L70
5 SEA ABB=ON PLU=ON (L71 OR L72)
L72
L73
L74
              1 SEA ABB=ON PLU=ON L29 AND (L68 OR L69 OR L70)
     FILE 'REGISTRY' ENTERED AT 10:20:26 ON 21 FEB 2008
     FILE 'ZCAPLUS' ENTERED AT 10:20:39 ON 21 FEB 2008
                 D STAT QUE L32
     FILE 'REGISTRY' ENTERED AT 10:20:59 ON 21 FEB 2008
```

FILE 'ZCAPLUS' ENTERED AT 10:21:01 ON 21 FEB 2008

D STAT QUE L73

L75 5 SEA ABB=ON PLU=ON (L73 OR L74)

FILE 'MEDLINE, EMBASE, BIOSIS' ENTERED AT 10:21:26 ON 21 FEB 2008 L76 1 SEA ABB=ON PLU=ON L73

FILE 'WPIX' ENTERED AT 10:21:40 ON 21 FEB 2008 77 2 SEA ABB=ON PLU=ON (L71 OR L72)

FILE 'STNGUIDE' ENTERED AT 10:21:48 ON 21 FEB 2008

FILE 'ZCAPLUS, EMBASE, WPIX' ENTERED AT 10:22:04 ON 21 FEB 2008 L78 5 DUP REM L75 L76 L77 (3 DUPLICATES REMOVED) ANSWERS '1-5' FROM FILE ZCAPLUS D IBIB ABS HITIND HITSTR L78 1-5 FILE 'REGISTRY' ENTERED AT 10:22:51 ON 21 FEB 2008

FILE 'ZCAPLUS' ENTERED AT 10:22:54 ON 21 FEB 2008

D STAT QUE L32 D STAT QUE L37

D STAT QUE L45 D STAT OUE L55

D STAT QUE L55 D STAT QUE L67

L79 17 SEA ABB=ON PLU=ON (L32 OR L37 OR L45 OR L55 OR L67) NOT (L73 OR L74)

D IBIB ABS HITIND HITSTR L79 1-17

FILE 'REGISTRY' ENTERED AT 10:26:43 ON 21 FEB 2008

FILE 'ZCAPLUS' ENTERED AT 10:26:46 ON 21 FEB 2008

D STAT QUE L64 D STAT QUE L65

L80 32 SEA ABB=ON PLU=ON (L64 OR L65) NOT (L79 OR L73 OR L74)
D IBIB ABS HITIND HITSTR L80 1-32

FILE HOME

FILE ZCAPLUS

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STRUCTURE FILE UPDATES: 20 FEB 2008 HIGHEST RN 1004854-20-9 DICTIONARY FILE UPDATES: 20 FEB 2008 HIGHEST RN 1004854-20-9

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FILE STNGUIDE

FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: Feb 15, 2008 (20080215/UP).

FILE MEDLINE

FILE LAST UPDATED: 20 Feb 2008 (20080220/UP). FILE COVERS 1949 TO DATE.

MEDLINE has been updated with the National Library of Medicine's revised 2008 MeSH terms. See HELP RLOAD for details.

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FILE EMBASE

FILE COVERS 1974 TO 20 Feb 2008 (20080220/ED)

 ${\tt EMBASE}$ is now updated daily. SDI frequency remains weekly (default) and biweekly.

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FILE BIOSIS

FILE COVERS 1926 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1926 TO DATE.

RECORDS LAST ADDED: 20 February 2008 (20080220/ED)

BIOSIS has been augmented with 1.8 million archival records from 1926 through 1968. These records have been re-indexed to match current BIOSIS indexing.

FILE WPIX

FILE LAST UPDATED: 20 FEB 2008 <20080220/UP>
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